

# STIC Search Report Biotech-Chem Library

## STIC Database Tracking Number: 142604

TO: Celine Qian

Location: REM-2A64/2C70

Art Unit: 1636

Wednesday, January 19, 2005

Case Serial Number: 10/009579

From: Edward Hart

**Location: Biotech-Chem Library** 

**REM-1A55** 

Phone: 571-272-2512

edward.hart@uspto.gov

## Search Notes

Examiner Qian,

Here are the results of the search you requested.

Please feel free to contact me if you have any questions.

**Edward Hart** 



## **WEST Search History**

Hide Items Restore Clear Cancel

DATE: Wednesday, January 19, 2005

Hide?	<u>Set</u> Name	Query	<u>Hit</u> Count			
DB=PGPB,USPT,JPAB,DWPI; PLUR=YES; OP=ADJ						
	L15	L8 and L14	4			
	L14	L13 same (promoter or regulat\$ or UTR or enhancer)	16			
	L13	human pancarcinoma associated epithelial glycoprotein-2 or EGP-2 or 17-1A or Ep-CAM	519			
	L12	human pancarcinoma associated epithelial glycoprotein-2 or EGP-2 17-1A or Ep-CAM	95			
	L11	L8 near5 (promoter or regulat\$ or UTR or enhancer)	67			
	L10	L8 same (promoter or regulat\$ or UTR or enhancer)	145			
	L9	L8 and (promoter or regulat\$ or UTR or enhancer)	1006			
	L8	carcinoma near3 (specific or select\$ or restrict\$)	1553			
DB=PGPB,USPT,USOC,JPAB,DWPI; PLUR=YES; OP=ADJ						
	L7	L5 same lung carcinoma	0			
2	L6	L5 and lung carcinoma	18			
	L5	L4 same (promoter or regulat\$ region or regulat\$ element or 5 prime UTR)	87			
	L4	carcinoma near3 (select\$ or specific\$ or prefer\$)	1683			
	L3	L1 near3 (promoter or regulat\$ region or regulat\$ element or 5 prime UTR)	1			
	L2	L1 and (promoter or regulat\$ region or regulat\$ element or 5 prime UTR)	307			
	L1	EGP-2 or Ep-CAM or 17-1A or GA733-2	573			

END OF SEARCH HISTORY

ATTN: Ed Hart

# 142604 SEARCH REQUEST FORM

Access DB#	
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# Scientific and Technical Information Center

CRFER

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Requester's Full Name: O Art Unit: 1636 Pho Mail Box and Bldg/Room Loc	eline Qi an (2010)  one Number 30 2 - 077  ation: 2464 R	Examiner #: 78770 Date: 1/13/04  7 Serial Number: 10/009, 579  csults Format Preferred (circle): PAPER DISK E-MAI	L
If more than one search is s	ubmitted, please prior ********	itize searches in order of need. **********************************	* *
Include the elected species or structi	ires, keywords, synonyms, ac terms that may have a special	ibe as specifically as possible the subject matter to be searched, cronyms, and registry numbers, and combine with the concept or meaning. Give examples or relevant citations, authors, etc., if and abstract.	
Title of Invention: Non-S	Squamous Lp.	thelium-Specific Transcription LEI] et al	<u>_</u>
Earlies: Priority Filing Date:	3/1/2000		
		on (parent, child, divisional, or issued patent numbers) along with the	
Please search	SEQIDNO:5	From 3115 bp - 3560 bp. (778 to -442 Figure 1)	
		₽. <del></del>	-
STAFF USE ONLY Storcher:	Type of Search  NA Sequence (#)/	Vendors and cost where applicable	
Searcher Phone #:	AA Sequence (#)	STN	
Date Searcher Picked Up://g/(///	Structure (#) Bibliographic	Questel/Orbit	
Date Completed: 1905	Litigation	Dr.Link	
Searcher Prem - Review Time	Fulltext	Sequence Systems	
Clerical Prep Time:	Patent Family		
FTO-1590 (8-01)	Other	Other (specify)	

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                                                                                                                                     YOU HAVE REQUESTED DATA FROM 37 ANSWERS - CONTINUE? Y/(N):y
                                                                                                                                     L3 ANSWER 1 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
                                                                                                                                           2004:1020014 CAPLUS
Welcome to STN International Enter x:x
                                                                                                                                      DN 142:5477
                                                                                                                                     TI Recombinant virus expressing an intact anti-tumor antibody containing
LOGINID:ssspta1633cxq
                                                                                                                                         human immunoglobulin constant regions and the therapeutic use thereof
                                                                                                                                     IN Qian, Qijun; Yang, Qin
PA Sino-Gene Biotechnology Ltd., Peop. Rep. China
 PASSWORD:
 TERMINAL (ENTER 1, 2, 3, OR 7):2
                                                                                                                                     SO PCT Int. Appl., 50 pp.
                                                                                                                                         CODEN: PIXXD2
 ******* Welcome to STN International
                                                                                                                                     DT Patent
                                                                                                                                     LA Chinese
FAN.CNT 1
                      Web Page URLs for STN Seminar Schedule - N. America "Ask CAS" for self-help around the clock
 NEWS 1
                                                                                                                                         PATENT NO.
                                                                                                                                                                    KIND DATE
                                                                                                                                                                                             APPLICATION NO.
 NEWS 2
  NEWS 3 SEP 01 New pricing for the Save Answers for SciFinder Wizard within
                                                                                                                                                                         A1 20041125 WO 2004-CN430
                                                                                                                                     PI WO 2004101777
 STN Express with Discover!
NEWS 4 OCT 28 KOREAPAT now available on STN
                                                                                                                                             W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
 NEWS 5 NOV 30 PHAR reloaded with additional data
NEWS 6 DEC 01 LISA now available on STN
NEWS 7 DEC 09 12 databases to be removed from STN on December 31, 2004
                                                                                                                                             GE, GH, GM, HK, HU, ID, IL, IN, IS, JP, KE, KG, KP, KK, Z, LC, LK, LR, LS, LT, LU, Y, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SL, SY, TB, EE, LE, CT, GE, CO, CM, CA, GN, CO, CW, MM, MB, NE
 NEWS 8 DEC 15 MEDLINE update schedule for December 2004
NEWS 9 DEC 17 ELCOM reloaded; updating to resume; current-awareness
 alerts (SDIs) affected

NEWS 10 DEC 17 COMPUAB reloaded; updating to resume; current-awareness alerts (SDIs) affected
 NEWS 11 DEC 17 SOLIDSTATE reloaded; updating to resume; current-
                                                                                                                                                SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
 awareness
                                                                                                                                     PRAI CN 2003-116733
                alerts (SDIs) affected
                                                                                                                                                                           A 20030430
 NEWS 12 DEC 17 CERAB reloaded; updating to resume; current-awareness alerts (SDIs) affected
                                                                                                                                     AB The present invention provides a recombinant virus comprising a chimeric gene which encodes an intact anti-tumor antibody contg. human lg const.
 NEWS 13 DEC 17 THREE NEW FIELDS ADDED TO IFIPAT/IFIUDB/IFICDB
                                                                                                                                          regions and the therapeutic use thereof. By inserting into the genome of
 NEWS 14 DEC 30 EPFULL: New patent full text database to be available on STN NEWS 15 DEC 30 CAPLUS - PATENT COVERAGE EXPANDED
                                                                                                                                         a recombinant virus a nucleotides acid sequence which simultaneously comprises. The cDNA sequences of both light chain and heavy chain gene of
                                                                                                                                         an intact anti-tumor antibody with human Ig const. regions are inserted into the genome of a recombinant virus, and the intact anti-tumor antibody
 NEWS 16 JAN 03 No connect-hour charges in EPFULL during January and
 February 2005
NEWS 17 JAN 11 CA/CAPLUS - Expanded patent coverage to include Russia
                                                                                                                                         can be expressed in tumor cells with high efficiency, thereby inhibit the
                                                                                                                                         growth and metastasis of tumors. In particular embodiments, the cDNA expressing human anti-EGFR antibody, or humanized antibody specific to
                (Federal Institute of Industrial Property)
 NEWS EXPRESS JANUARY 10 CURRENT WINDOWS VERSION IS V7.01a,
                                                                                                                                         human Her2, and chimeric human-mouse anti-CD20 antibody are prepd. and
                                                                                                                                         inserted into a replication deficient adenovirus. The anti-tumor activity of chimeric human-mouse anti-CD20 antibody is tested in breast cancer cell
             MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
             AND CURRENT DISCOVER FILE IS DATED 10 JANUARY 2005
                                                                                                                                         line BT-474 and a nude mouse implemented with breast cancer cell line
                                                                                                                                          SK-OV-3.
 NEWS HOURS STN Operating Hours Plus Help Desk Availability
NEWS INTER
NEWS LOGIN Welcome Banner and News Items
NEWS PHONE Direct Dial and Telecommunication Network Access to STN
                                                                                                                                     RE.CNT 4
                                                                                                                                                        THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
                                                                                                                                     RECORD
                                                                                                                                                 ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                                                                                                                     L3 ANSWER 2 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2004:927015 CAPLUS
 NEWS WWW
                         CAS World Wide Web Site (general information)
                                                                                                                                     DN 141:394059
 Enter NEWS followed by the item number or name to see news on that
                                                                                                                                     TI Human EpCAM or TAg-25, fragments, chimeric derivatives, antibodies and
                                                                                                                                     numan ≝p∟AM or IAg.-25, tragments, chimenc derivatives, antibodic conjugates for cancer diagnosis and therapy
IN Punnonen, Juha; Apt, Doris; Neighbors, Margaret; Leong, Steven R. PA Maxygen, Inc., USA
SO PCT Int. Appl., 273 pp.
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                                                                                                                                         CODEN: PIXXD2
                                                                                                                                     DT Patent
                                                                                                                                     LA English
  FAN.CNT 1
                                                                                                                                         PATENT NO.
                                                                                                                                            NO 2004093808 A2 20041104 WO 2004-US12280 20040419
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG US 2003-464780P P 20030422
                                                                                                                                                                    KIND DATE
                                                                                                                                                                                             APPLICATION NO.
 FILE 'HOME' ENTERED AT 16:15:29 ON 19 JAN 2005
                                                                                                                                     PI WO 2004093808
 => FIL BIOSIS EMBASE CAPLUS
COST IN U.S. DOLLARS
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PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
                                                                                                                                     PRAI US 2003-464780P
                                                                                                                                                                         P 20030422
                                                                                                                                     AB The invention provides novel polypeptides, including novel tumor-assocd. antigens, and related nucleic acids, vectors, cells, fusion nucleic acids
COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)
                                                                                                                                         or polypeptides, ligands and antibodies. The invention also provides compns. comprising such polypeptides, nucleic acids, vectors, cells, and
=> s EGF 2 or epithelial glycoprotein 2 or Ep CAM or 17 1A or GA733 2
L1 1526 EGF 2 OR EPITHELIAL GLYCOPROTEIN 2 OR EP CAM OR 17 1A
                                                                                                                                          antibodies, and methods of producing and using the same.
                                                                                                                                     L3 ANSWER 3 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2004:902213 CAPLUS
OR GA733 2
=> s I1 and (promoter or regula? element or regulat? region or 5 UTR)
L2 52 L1 AND (PROMOTER OR REGULA? ELEMENT OR REGULAT?
REGION OR 5
                                                                                                                                     DN 141:378849
                                                                                                                                     TI Immunogenic recombinant antibodies for use as vaccines against infection,
                                                                                                                                         autoimmune disease and cancer in primate such as human
Loibner, Hans; Himmler, Gottfried; Waxenecker, Guenter, Schuster, Manfred;
             UTR)
                                                                                                                                     PA Igeneon Krebs-Immuntherapie Forschungs- und Entwicklungs-A.-G., Austria SO PCT Int. Appl., 59 pp.
=> dup rem 12
PROCESSING COMPLETED FOR L2
            37 DUP REM L2 (15 DUPLICATES REMOVED)
                                                                                                                                         CODEN: PIXXD2
                                                                                                                                     DT Patent
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DATE

DATE

20040429

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A English
  FAN CNT 1
         PATENT NO
                                                     KIND DATE
                                                                                               APPLICATION NO.
  PI WO 2004091655
                                                              A2 20041028 WO 2004-EP4059
                                                                                                                                                             20040416
             NO 2004091655

A2 20041028 WO 2004-EP4059

20040416
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                     TD. TG
  PRAI AT 2003-599
                                                                 20030417
 AB The invention refers to an immunogenic recombinant antibody designed for immunization of primates comprising at least a part of a murine IgG2a subtype amino acid sequence and a mammalian glycosylation. The antibody
        is a chimeric, humanized, monoclonal, anti-idiotypic, or bi-isotopic antibody or fragment. The antigen is an epitope or mimotope of tumor-assocd. antigen, epithelial cell adhesion mol., Lewis Y antigen, NCAM, CEA, T cell epitope, carbohydrate, sialyi-Tn, Globo-H, glycolipid,
         GD2, GD3 or GM2.
 L3 ANSWER 4 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
  AN 2004:634026 CAPLUS
 DN 141:172878
 TI Engineering of glycosylation profile of antibody Fc region to increase Fc
 receptor binding affinity and effector function for treating cancer IN Umana, Pablo; Bruenker, Peter; Ferrara, Claudia; Suter, Tobias
            Glycart Biotechnology Ag, Switz.
       PCT Int. Appl., 231 pp.
CODEN: PIXXD2
 DT Patent
LA English
 FAN.CNT 1
        PATENT NO.
                                                     KIND DATE
                                                                                               APPLICATION NO.
                                                                                                                                                        DATE
 PI WO 2004065540
                                                            A2 20040805 WO 2004-IB844
                                                                                                                                                          20040122
             NO 200401824 20040185 WO 2004-IB844 20040122
W: AE, AE, AG, AL, AL, AM, AM, AM, AT, AT, AU, AZ, AZ, BA, BB, BG,
BG, BR, BR, BW, BY, BY, BZ, BZ, CA, CH, CN, CN, CO, CO, CR, CR,
CU, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EC, EC, EE, EE, EG, ES,
ES, FI, FI, GB, GD, GE, GE, GH, GM, HR, HR, HU, HU, IDI, IN, II, IN,
IS, JP, JP, KE, KE, KG, KG, KP, KP, KP, KR, KZ, KZ, KZ, LC,
LK, LR, LS, LS, LT, LU, LV, MA, MD, MD, MG, MK, MN, MW, MX, MX,
MZ, MZ, NA, NI
S 2004241817 A1 20041202 US 2004-761435 20042182
MZ, MZ, NA, NI
US 2004241817 A1 20041202 US 2004-761435
PRAI US 2003-441307P P 20030122
US 2003-491254P P 20030731
US 2003-495142P P 20030815
US 2003-495142P P 20030815

AB The present invention relates to nucleic acid mols., including fusion constructs, having catalytic activity and the use of same in glycosylation engineering of host cells to generate polypeptides with improved therapeutic properties, including antibodies with increased For receptor binding and increased effector function. The engineered proteins or antibodies comprise Golgi localization domain of Golgi resident polypeptide such as .beta.(1,4)-N-acetylglucosaminyltransferase III,
        .beta.(1,4)-galactosyltransferase, mannosidase II, .beta.(1,2)-N-acetylglucosaminyltransferase I, .beta.(1,2)-N-acetylglucosaminyltransferase II, mannosidase I, .alpha.-mannosidase II,
       and .alpha.1-6 core fucosyltransferase. The effector function includes Fc-mediated cellular cytotoxicity of NK cells, macrophage,
        polymorphonuclear cells and monocytes; signaling of apoptosis induction; maturation of dendritic cells; or T cell priming. The engineered antibodies include antibodies or humanized antibodies specific to human
       neuroblastoma, renal cell carcinoma, colon carcinoma, breast carcinoma, lung carcinoma, ***17*** - ***1A*** antigen, CD20, CD22, CD30, CD40, PSMA, EGFR, PSCA, HLA-DR, MUC1, EpCAM, etc.
L3 ANSWER 5 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
          2004:80710 CAPLUS
DN 140-144706
 TI Production of recombinant antibodies comprising one common light chain and
three different heavy chains for diagnosis and therapy
IN Van Berkel, Patrick Hendrikus Cornelis; Brus, Ronald Hendrik Peter; Bout,
        Abraham; Logtenberg, Ton
Crucell Holland B.V., Neth.
SO PCT Int. Appl., 186 pp.
CODEN: PIXXD2
 DT Patent
   A English
FAN CNT 1
       PATENT NO.
                                                     KIND DATE
                                                                                              APPLICATION NO.
                                                                                                                                                       DATE
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A2 20040129 WO 2003-EP7690 A3 20041104

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,

20030715

PI WO 2004009618

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TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
EP 2002-77953 A 20020718
S 2002-397086P P 20020718
O 2003-EP50201 A 20030527
The invertice provides methods for producing mixts of antihodies from a
  PRAI EP 2002-77953
US 2002-397066P
         WO 2003-EP50201
  AB The invention provides methods for producing mixts. of antibodies from a
         single host cell clone. Thereto a nucleic acid sequence encoding a light chain, and nucleic acid sequences encoding different heavy chains are
       chain, and nucleic acid sequences encoding different neary chains are expressed in a recombinant host cell. The antibodies in the mixts. according to the invention suitably comprise identical light chains paired to different heavy chains capable of pairing to the light chain, thereby forming functional antigen binding domains. Antibodies exemplified in the invention include VL and VH of clones K53 (against CD46), UBS-54 (against ***Ep**** - ****CAM**** ), 02-237 (against CD45), B28 (against CD22), II-2 (against CD72) and I-2 (against HLA-DR class II). Such mixts. can be used in a variety of fields.
  L3 ANSWER 6 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation.
 on
         STN
                                                                                              DUPLICATE 1
  AN 2004:441672 BIOSIS
  DN PREV200400446570
         Use of the EGP-2/ ***Ep*** - ***CAM*** ***promoter*** for targeted expression of heterologous genes in carcinoma derived cell lines.
  AU McLaughlin, Pamela M. J.; Tizpis, Monika; Kroesen, Bart-Jan; Helfrich, Wijnand; Terpstra, Peter, Dokter, Wim H. A.; Ruiters, Marcel H. J.; de Leij, Lou F. M. H.; Harmsen, Martin C. [Reprint Author]
 CS Dept Pathol and Lab MedSect Med Biol, Univ Groningen Hosp, Hanzepl 1, NL-9713 GZ, Groningen, Netherlands
         m.c.harmsen@med.rug.nl
 SO Cancer Gene Therapy, (September 2004) Vol. 11, No. 9, pp. 603-612. print. ISSN: 0929-1903 (ISSN print).
  DT Article
 LA English
ED Entered STN: 17 Nov 2004
 Last Updated on STN: 17 Nov 2004
AB EGP-2, also known as ***Ep*** - ***CAM****, is expressed at high
         levels on the surface of most carcinomas and is therefore considered an
         attractive target for anticancer strategies. To explore the mechanisms regulating the expression of EGP-2, sequences 3.4 kb upstream of the
         transcription start site were isolated and assayed for their ability to control the expression of the EGP-2 cDNA, the green fluorescent protein, the luciferase reporter gene and the thymidine kinase and cytosine
         deaminase suicide genes. Expression of these chimeric constructs as assessed in a range of different cell lines was restricted to cell lines expressing EGP-2. In addition, only cells expressing EGP-2 were sensitive
       expressing EGP-2. In addition, only cells expressing EGP-2 were sensity for gancyclovir after being transiently transfected with EGP-2

***promoter*** -driven thymidine kinase. Deletion analyses defined 687 bp upstream as the basic proximal

***promoter*** region, which could confer epithelial-specific expression to the GFP reporter gene in vitro. As these EGP-2 sequences can confer

***promoter*** activity to reporter and suicide genes in an EGP-2 restricted manner, they may be useful for gene therapy of EGP-2 expressing carcinomas.
L3 ANSWER 7 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003-472621 CAPLUS
           139:51600
         Chimeric antigen comprising CD36-binding domain for enhancing vaccine
         immune response
          Cox, William I.; Alexander, Jeannine P.; Goebel, Scott
            Aventis Pasteur Limited, Can.
 SO PCT Int. Appl., 54 pp. CODEN: PIXXD2
 DT Patent
  LA English
FAN.CNT 1
        PATENT NO.
                                                        KIND DATE
                                                                                                     APPLICATION NO
PI WO 2003050268
PI WO 2003050268 A2 20030619 WO 2002-US39885 20021212 WO 2003050268 A3 20040708

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MX, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, BB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

US 2004241652 A1 20041202 US 2002-317821 20021212

PRAI US 2001-341771P P 20011212

AB The invention relates to reagents and methods for enhancing an immune
                                                                 A2 20030619 WO 2002-US39885
                                                                                                                                                                         20021212
  AB The invention relates to reagents and methods for enhancing an immune
       response using CD36 binding region/antigen hybrid polypeptides or polynucleotides encoding the hybrid polypeptides. The antigen is gp100, MART/Melan A, gp75/TRP-1, tyrosinase, NY-ESO-1, melanoma proteoglycan, MAGE-1, MAGE-2, MAGE-3, MAGE-4, MAGE-6, MAGE-12, BAGE, GAGE-1,
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RAGE, N-actylglucosaminyltransferase V, p15, .beta.-catenin, MUM-1, cyclin dependent kinase 4, p21 ras, BCR-abl, p53, p185 HER2/neu, EGF receptor,

CEA antigen, MUC-1, EBNA-1, E7, E6, prostate-specific antigen, prostate specific membrane antigen, KSA, or NY-BR-1. L3 ANSWER 8 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003:472615 CAPLUS DN 139:30800 TI Streptavidin expressed gene fusions with single-chain antibodies and their use as targeting vehicles for diagnosis and treatment of cancer IN Goshorn, Stephen Charles; Graves, Scott Stoll; Schultz, Joanne Elaine; Lin, Yukang; Sanderson, James Allen; Reno, John M.; Dearstyne, Erica A. PA Neorx Corporation. USA SO PCT Int. Appl., 156 pp. CODEN: PIXXD2 DT Patent LA English FAN.CNT 5 PATENT NO. KIND DATE APPLICATION NO. WO 2003050260
A2 20030619
WO 2003050260
A3 20041125
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PI, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
US 2003103948
A1 20030605
US 2002-150762
20020517
US 20031143233
A1 20030731
US 2002-244821
20020916
RAI US 2001-13173
A 20011207 PI WO 2003050260 A2 20030619 WO 2002-US39429 20021206 20011207 20020517 Α US 2002-244821 20020916 US 1999-137900P US 1999-168976P 19990607 19991203 US 2000-589870 A2 20000605 from these vectors are provided. In particular embodiments, fusion proteins comprising a single-chain antibody and genomic streptavidin are provided as are vectors encoding the same. The single-chain antibodies are directed to cell surface antigens, or cell-assocd. stromal or matrix antigens, including, but not limited to, CD20, CD22, CD25, CD45, CD52, CD56, CD57, EGP40 (or EPCAM or KSA), N-CAM, CEA, TAG-72, .gamma.transferase, mucins (MUC1 through MUC7), human .beta.-chorionic gonadotropin, EGF receptor, interleukin-2 receptor, her2/neu, Lewis Y, gangliosides GD2 and GM2, tenascin, sialylated tenascin, somatostatin, activated tumor stromal antigen, or recoangiogenic antigens. Generically, a single-chain Fv/streptavidin (scFvSA) fusion protein is expressed from the genetic fusion of the single-chain antibody of the variable regions to the genomic streptavidin of Streptomyces avidinii. The scFv gene consists of the variable regions of the light and heavy chains sepd. by a DNA linker sequence. The streptavidin coding sequence is joined to the 3'-terminus of the scFv gene, and the two genes are sepd, in-frame by a second DNA linker sequence. The signal sequence from the streptavidin gene is fused at the 5'-terminus of the scFvSA gene to direct expression gene is lased at the 3-terminals of the scrvsA gene to direct expression to the Escherichia coli periplasmic space. The scrvSA gene is under control of the lac \*\*\*promoter\*\*\*, and the expressed fusion protein is extd. and purified from E. coli and forms a sol. tetramer of apprx.173,000 mol. wt. Also provided, are methods of using the fusion proteins of the present invention, in the absence and presence of a radiation-sensitizing agent (e.g., Gemcitabine), and in particular, the use of scFvSA fusion proteins as diagnostic markers or as cell-specific targeting agents. Immunivest Corporation, USA

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PRAI US 2001-13173
US 2002-150762
AB The present invention provides vectors for expressing genomic streptavidin fusion cassettes. In the various embodiments, fusion proteins produced
glutamyl
L3 ANSWER 9 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
AN 2003:335307 CAPLUS
DN 138:350812
TI Use of nucleic acid and protein profiling and histology of fixed cells in
a single sample in the early diagnosis of disease
IN O'Hara, Shawn Mark; Zweitzig, Daniel; Foulk, Brad
SO PCT Int. Appl., 105 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 2
     PATENT NO.
                                          KIND DATE
                                                                             APPLICATION NO
                                                                                                                             DATE
                                               A2 20030501 WO 2002-US34570
A3 20040108
PI WO 2003035895
                                                                                                                                  20021028
     WO 2003035895
         /O 2003035895 A3 20040108

W. AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
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FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
EP 1438419 A2 20040721 EP 2002-795565 20021028
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK
PRAI US 2001-330669P P 20011028
US 2002-369945P P 20020404
WO 2002-US34570 W 20021028
AB A highly sensitive assay is disclosed which utilizes a method for gene specific primed amplification of mRNA libraries from rare cells and rare transcripts found in blood. The assay allows detection of rare, mRNA (10 copies/cell) found in 1 to 10 cells isolated through immunomagnetic
       copies/cell) found in 1 to 10 cells isolated through immunomagnetic enrichment. The assay is an improvement over multiplex PCR and allows efficient detection of rare coding sequences for circulating carcinoma
       cells in the blood. The methods are useful in profiling of cells isolated from tissues or body fluids and serves as an adjunct to clin. diagnosis of
       diverse carcinomas including early stage detection and classification of
      circulating tumor cells. Monitoring of nucleic acid and protein profiles of cells either in conventional or microarray formats, facilitates
       management of therapeutic intervention including staging, monitoring
       response to therapy, confirmation of remission and detection of
       regression
 L3 ANSWER 10 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
  AN 2003:590597 CAPLUS
 DN 139:144951
       Preparation of fusion genes encoding streptavidin and single chain
       antibody and methods of therapeutic use thereof
      Goshorn, Stephen Charles; Graves, Scott Stoll; Schultz, Joanne Elaine;
Lin, Yukang; Sanderson, James Allen; Reno, John M.; Dearstyne, Erica A.
        NeoRx Corporation, USA
 SO U.S. Pat. Appl. Publ., 89 pp., Cont.-in-part of U.S. Ser. No. 150,762. CODEN: USXXCO
 DT Patent
LA English
      PATENT NO.
                                          KIND DATE
                                                                           APPLICATION NO.
                                                                                                                       DATE
 PI US 2003143233
                                                       20030731
                                                                              US 2002-244821
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      US 2003095977
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       US 2003103948
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                                                      20030605
                                                                            US 2002-150762
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      WO 2003050260
WO 2003050260
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A3
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20041125
                                                                             WO 2002-US39429
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WO 2003050260 A3 20041125
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-137900 P 19999607
US 1999-168976P P 19991203
US 2000-589870 A2 20000605
US 2001-13173. A2 200011207
       US 2001-13173
                                           A2 20011207
      US 2002-150762
US 2002-244821
                                            A2 20020517
A 20020916
 AB The present invention provides vectors for expressing genomic streptavidin
      fusion cassettes and therapeutic uses. In the various embodiments, fusion proteins produced from these vectors are provided. In particular embodiments, fusion proteins comprising a single chain antibody and
      genomic streptavidin are provided as are vectors encoding the same. Also provided, are methods of using the fusion proteins of the present
      invention, in the absence and presence of a radiation-sensitizing agent, and in particular, the use of scFvSA fusion proteins as diagnostic markers
      or as a cell specific targeting agents.
L3 ANSWER 11 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003;435061 CAPLUS DN 139:21033
       Vectors expressing soluble form of single chain antibody and streptavidin
       (scFvSA) fusions and uses thereof as diagnostic markers or as cell
       specific targeting agents
 IN Goshorn, Stephen Charles; Graves, Scott Stoll; Schultz, Joanne Elaine;
Lin, Yukang; Sanderson, James Allen; Reno, John M.; Dearstyne, Erica A.
PA NeoRx Corporation, USA
      U.S. Pat. Appl. Publ., 84 pp., Cont.-in-part of U.S. Ser. No. 13,173. CODEN: USXXCO
 DT Patent
LA English
FAN.CNT 5
      PATENT NO.
                                          KIND DATE
                                                                           APPLICATION NO.
                                                                                                                       DATE
 PI US 2003103948
                                                       20030605
                                                                              US 2002-150762
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      US 2003095977
US 2003143233
                                                     20030522 US 2001-13173
20030731 US 2002-244821
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20020916
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      WO 2003050260
                                                      20030619
                                                                             WO 2002-US39429
                                                                                                                          20021206
          O 2003050260 A3 20041125
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
      WO 2003050260
              CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
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LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, ŜL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-137900P P 19991203
US 1999-188976P P 19991203
US 2000-589870 A2 2000-0615
        US 2000-589870
US 2001-13173
                                                   A2 20000605
A2 20011207
                                                     A2 20020517
         US 2002-150762
         US 2002-244821
                                                             20020916
   AB The present invention provides vectors for expressing Streptomyces
         avidinii genomic streptavidin (SA) fusion cassettes. A genomic
        streptavidin expressed gene fusion is expressed as a sol. protein into the periplasmic space of bacteria and undergoes spontaneous folding. Such
        expression offers the advantage that the periplasm is a low biotin environment and one need not purify and refold the protein under harsh denaturing conditions that may prove fatal to the polypeptide encoded by a
       denaturing conditions that may prove tatal to the polypeptide encoded by a 
heterologous nucleic acid mol. fused to the genomic streptavidin nucleic 
acid mol. In the various embodiments, fusion proteins produced from these 
vectors are provided. In particular embodiments, fusion proteins 
comprising a single chain antibody and streptavidin (scFvSA) are provided 
as are vectors encoding the same. The single chain antibodies are 
directed to cell surface antigens or cell-assocd. stromal or matrix 
proteins such as CD20, CD45, CD22, CD52, CD56, CD57, EGP40, NCAM,
  CEA,
        TAG-72, mucins (MUC1-7), 13HCG, EGF receptor, IL-2 receptor, her2/neu, Lewis Y, GD2, GM2, tenascin, sialylated tenascin, somatostatin, activated tumor stromal antigen or neoangiogenic antigens. Also provided, are
        methods of using the fusion proteins of the present invention, in the absence and presence of a radiation-sensitizing agent, and in particular,
         the use of scFvSA fusion proteins as diagnostic markers or as a cell
         specific targeting agents.
 L3 ANSWER 12 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003;396269 CAPLUS
  DN 138:400405
  TI Streptavidin-antibody fusion proteins for diagnosis and specific cell
        targeting
  IN Goshorn, Stephen Charles; Graves, Scott Stoll; Schultz, Joanne Elaine;
  Lin, Yukang, Sanderson, James Allen, Reno, John M. PA Neorx Corporation, USA
 SO U.S. Pat. Appl. Publ., 72 pp., Cont.-in-part of U.S. Ser. No. 589,870 CODEN: USXXCO
  DT Patent
 LA English
FAN.CNT 5
       PATENT NO.
                                                  KIND DATE
                                                                                        APPLICATION NO.
                                                                                                                                             DATE
  PI US 2003095977
                                                                 20030522 US 2001-13173
                                                                20030605 US 2002-150762
20030731 US 2002-244821
         US 2003103948
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         US 2003143233
         WO 2003050260
                                                                 20030619 WO 2002-US39429
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WO 2003050260 A3 20041125
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-168976P
P 19991203
US 2000-589870
A2 20000605
        WO 2003050260
                                                       A3 20041125
        US 2000-589870
US 2001-13173
                                                   A2 20000605
A2 20011207
                                                             20020517
20020916
         US 2002-150762
                                                    A2
        US 2002-244821
       fusion cassettes and fusion protein produced from the vectors. In particular embodiments, fusion proteins comprising a single chain antibody
         and genomic streptavidin are provided as are vectors encoding the same.
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AB The present invention provides vectors for expressing genomic streptavidin Also provided are methods of using the fusion proteins of the present invention, and in particular, the use of scFvSA fusion proteins as diagnostic markers or as a cell specific targeting agents. The single chain antibodies are directed to cell surface antigens or cell-assood. stromal or matrix protein such as CD20, CD45, CD22, CD52, CD56, CD57, CD60, CD67, CD67, CD60, CD67, CD67, CD60, CD67, CD67, CD60, CD67, CD60, CD67, CD60, CD67, CD60, CD67, CD60, CD67, CD67, CD60, CD67, CD60, CD67, CD60, CD67, EGP40, NCAM, CEA, TAG-72, mucins (MUC1-7), 13HCG, EGF receptor, IL-2 receptor, her2/neu, Lewis Y, GD2, GM2, tenascin, sialylated tenascin, somatostatin, activated tumor stromal antigen or neoangiogenic antigens.

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L3 ANSWER 13 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003:704297 CAPLUS
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DN 139:346666

TI Cloning and characterisation of a 1.1kb fragment of the carcinoma-associated epithelial cell adhesion molecule \*\*\*promoter\*\*\*

AU Gires, Olivier, Eskofier, Sylvia; Lang, Stephan; Zeidler, Reinhard; Muenz,

CS Clinical Cooperation Group Molecular Oncology, GSF-Research Center for

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Health and Environment, and Department of Otorhinolaryngology,
       Ludwig-Maximilians-University, Munich, D-81377, Germany
O Anticancer Research (2003), 23(4), 3255-3261
CODEN: ANTRD4; ISSN: 0250-7005
  PB International Institute of Anticancer Research
          Journal
  LA English

    The epithelial cell adhesion mol. (EpCAM) is a transmembrane protein assocd, with a variety of carcinomas, where EpCAM is often strongly

  AB
       up-regulated or, as in the case of squamous cell carcinomas, de novo expressed. The mol. mechanisms underlying the transcriptional regulation
       of EpCAM are poorly understood. So far, a 570bp fragment has been cloned and shown to have specific transcriptional activity, which was neg.-regulated upon the induction of the transcription factor
       NF. vkappa.B. In the present study we have cloned a 1100bp fragment of the EpCAM ***promoter*** contg. the 570bp fragment and addnl. 550bp upstream. We demonstrate that both fragments have strong synergistic
       effects with respect to transcriptional activity in EpCAM-pos. cells. Furthermore, the 1100bp fragment was likewise neg.-regulated upon
 THF.alpha. and IFN.alpha. treatment, thus retaining silencer sequences.

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS
  RECORD
                  ALL CITATIONS AVAILABLE IN THE RE FORMAT
  L3 ANSWER 14 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
 AN 2002:575292 CAPLUS
DN 137:153381
  TI Genes overexpressed in prostate disorders as diagnostic and therapeutic
 targets
IN Hampton, Garret Malcolm; Welsh, John Barnard
PA IRM, LLC, Bermuda
SO PCT Int. Appl., 55 pp.
CODEN: PIXXD2
 DT Patent
LA English
       PATENT NO.
                                            KIND DATE
                                                                              APPLICATION NO.
                                                                                                                            DATE
AB Disclosed are methods for diagnosing, monitoring the progression of, and
       treating a prostate disorder based upon genes that are differentially expressed in prostate disorders. Also disclosed are methods for
      expressed in prostate disorders. Also disclosed are memods to identifying agents useful in the treatment of a prostate disorder, methods for monitoring the efficacy of a treatment for a prostate disorder, methods for inhibiting the proliferation of a prostate cell, and prostate-specific vectors including the ""promoter" of these genes. A dendrogram of 55 exptl. samples that are grouped according to overall similarity in level of expression of a subset of 3,530 genes that have varied meet agrees the samples in provided. Expression levels of highly
      varied most across the samples is provided. Expression levels of highly ranked genes in normal and malignant prostate tissues are provided. Furthermore, the top 25 or 50 genes (with ref. GenBank accession nos.) overexpressed in prostate malignant tissues or cell lines are identified as the diagnostic markers and therapeutic targets for prostate related
       disorders. They include genes for hepsin, prostate differentiation factor, alpha-methylacyl-CoA racemase, fatty acid synthase, and prostate
       specific antigen (alternative splice form 2 and 3). Specifically, the amplification of two marker genes (hepsin and PLAB) are detected at the mRNA level from selected prostate tissues.
  L3 ANSWER 15 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
         2002:10532 CAPLUS
  DN 136:84702
       Novel ligands for CD28 and CTLA-4 created by shuffling of mammalian B7-1
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iligand cDNAs with possible therapeutic use as co-stimulatory molecules
IN Punnonen, Juha; Lazetic, Alexandra L. L.; Leong, Steven R.; Chang,
Chia-Chun Jean; Apt, Doris; Gustafsson, Claes
PA Mavygen, Inc., USA
SO PCT Int. Appl., 364 pp.
        CODEN: PIXXD2
DT Patent
LA English
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APPLICATION NO.

DATE

KIND DATE

FAN.CNT 2 PATENT NO.

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CODEN: PIXXD2
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LA English
                                                                     A2 20020103 WO 2001-US19973
C2 20030206
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   PL WO 2002000717
           WO 2002000717

    I/O 2002000/17
    I/O 200200/17
    I/O 200200/17

           WO 2002000717
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         RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

CA 2411828 AA 20020103 CA 2001-2411828 20010622

EP 1360290 A2 20031112 EP 2001-952193 20010622

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR

JP 2004513878 T2 20040513 JP 2002-505839 20010622

US 2003138881 A1 20030724 LIS 2004 2004
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                                                                            20040513 JP 2002-505839
20030724 US 2001-32214
2 20040408 WO 2002-US19898
3 20041028
          US 2003138881
WO 2004029197
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WO 2004029197 A3 20041028
WO 2004029197 A3 20041028
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
EP 1497426 A2 20050119 EP 2002-807658 20020621
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
PRAI US 2000-241245P P 20001017
US 2001-888324 A2 200110822
WO 2001-US19898 W 20020621
AB The invention provides polynucleotides and polypeptides encoded therefrom
           WO 2004029197
                                                                                                                                                                                                                                                                                2001:338579 CAPLUS
                                                                                                                                                                                                                                                                                134:365705
  AB The invention provides polynucleotides and polypeptides encoded therefrom having advantageous properties, including an ability of the polypeptides
         naving advantageous properties, including an ability of the polypeptides to preferentially bind a CD28 or CTLA-4 receptor at a level greater or less than the ability of human B7-1 to bind CD28 or CTLA-4, or to induce or inhibit altered level of T cell proliferation response greater compared to that generated by human B7-1. The polypeptides and polynucleotides of the invention are useful in therapeutic and prophylactic treatment
                                                                                                                                                                                                                                                                   PA Maxygen, Inc., USA
SO PCT Int. Appl., 109 pp.
                                                                                                                                                                                                                                                                             CODEN: PIXXD2
                                                                                                                                                                                                                                                                    חד
                                                                                                                                                                                                                                                                               Patent
                                                                                                                                                                                                                                                                    LA English
          methods, gene therapy applications, and vaccines. Novel ligands were generated by shuffling of sequences from cDNAs for B7-1 ligands from human, rhesus monkey, baboon, orangutan, cow, cat and rabbit. Ligands
                                                                                                                                                                                                                                                                    FAN.CNT 1
                                                                                                                                                                                                                                                                           PATENT NO.
          were screened for using a FACS assay. CDNA libraries were introduced into animal cells that were then screened for their ability to bind a labeled
                                                                                                                                                                                                                                                                   PI WO 2001032712
WO 2001032712
                                                                                                                                                                                                                                                                                                                                       АЗ
          CD28 or CTLA-4 using FACS. Clones were screened for their preferential binding of CD28 vs. CTLA-4. Candidate clones were then tested for their ability to stimulate T cell proliferation.
   L3 ANSWER 16 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
   Corporation. on
                                                                                                      DUPLICATE 2
   AN 2002:523713 BIOSIS
   DN PREV200200523713
   TI Murine spermatogonial stem cells: Targeted transgene expression and
         purification in an active state.
   AU Giuili, Galicia; Tomljenovic, Andrea; Labrecque, Nathalie
           Oulad-Abdelghani, Mustapha, Rassoulzadegan, Minoo, Cuzin, Francois
          [Reprint author]
  CS Unite 470 de l'INSERM, Universite de Nice, F-06108, Nice Cedex 2, France
         fcuzin@unice.fr
  SO EMBO Reports, (August, 2002) Vol. 3, No. 8, pp. 753-759. print. ISSN: 1469-221X.
   DT Article
  LA English
  ED Entered STN: 9 Oct 2002
           Last Updated on STN: 9 Oct 2002
  AB A 400 bp fragment of the spermatogonia-specific Stra8 locus was sufficient to direct gene expression to the germinal stem cells in transgenic mice.
         A fractionation procedure was devised, based on immunomagnetic sorting of cells in which the ***promoter*** drives the expression of a surface
                                                                                                                                                                                                                                                                            agents are provided.
         cells in which the ""promoter" drives the expression of a surface functionally neutral protein tag. The purified cells expressed the known molecular markers of spermatogonia Rbm, cyclin A2 and ""EP" - ""Cam", and the beta1- and alpha6-integrins characteristic of the stem cell fraction. A 700-fold enrichment in stem cells was determined by
                                                                                                                                                                                                                                                                    Corporation, on
                                                                                                                                                                                                                                                                           STN
         the ability of the purified fractions to re-establish spermatogenesis in germ cell-depleted recipient testes.
 L3 ANSWER 17 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
  AN 2001:816487 CAPLUS
 DN 135:356752
IT Epitope synchronization in antigen presenting cells
IN Simard, John J. L.; Diamond, David C.; Lei, Xiang-Dong
PA CTL Immunotherapies Corp., USA
                                                                                                                                                                                                                                                                           [Reprint author]
  SO PCT Int. Appl., 131 pp.
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A2 20011108 WO 2001-US13806
                                                                                                                                                     20010427
                                                       A3 20020411
            W: AE, AG, AL, AM, AT, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EE, EE, ES, FI, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR,
                  KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK, SL, TJ, TM, TR,
                   TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
CA 2405363 AA 20011108 CA 2001-2405363 20010427
EP 1276896 A2 20030122 EP 2001-930922 20010427
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
JP 2003535824 T2 20031022 JP 2001-579836 20010427
PRAI US 2000-560455 A 20000428
US 2000-561074 'A 20000428
US 2000-561571 A 20000428
US 2000-561572 A 20000428
WO 2001-US13806 W 20010427
AB Disclosed herein are vaccines and methods for inducing an immune response against cancer cells and cells infected with intracellular parasites.
       against cancer cells and cells infected with intracellular parasites.
       Vaccines having housekeeping epitopes are disclosed. The housekeeping epitope is formed by housekeeping proteasomes in peripheral cells, but not
        by professional antigen presenting cells. A vaccine contg. a housekeeping
       epitope that is derived from an antigen assocd, with a peripheral target cell can thus direct an immune response against the target cell. Methods
        of treatment are also disclosed, which involve administering a vaccine
       having a housekeeping epitope.
 L3 ANSWER 18 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
        Antibody diversity generation
Karrer, Erik; Bass, Steven H.; Whalen, Robert; Patten, Phillip A.
                                                  KIND DATE
                                                                                         APPLICATION NO.
                                                                                                                                              DATE
                                                        A2 20010510 WO 2000-US30247
A3 20020321
                                                                                                                                                    20001101
            W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, RW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1230269 A2 20020614 EP 2000-976844 20001101

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

PRAI US 1999-163370P P 19991103

US 2000-176002P P 20000112

WO 2000-US30247 W 20001101

AB Methods for improving antibodies by a variety of DNA diversification and selection procedures are provided. Improvements include increases in
       selection procedures are provided. Improvements include increases in affinity, alterations in specificity and effector function, as well as reduced antigenicity, e.g. humanization. Libraries of recombinant
       antibody sequences are provided, as are cells expressing members of such libraries. Novel phage display vectors are provided. Methods for the coevolution of an antibody and its cognate antigen are provided.
       Coevolution is used to evolve HIV envelope proteins with increased
       antigenicity and broadly neutralizing antibodies that interact therewith.
       Methods of improving antibodies for use in the detection of biol. warfare
 L3 ANSWER 19 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
                                                                                 DUPLICATÉ 3
AN 2001:300365 BIOSIS
DN PREV200100300365
TI The ***epithelial*** ***glycoprotein*** ***2*** (EGP-2)
****promoter*** -driven epithelial-specific expression of EGP-2 in
       transgenic mice: A new model to study carcinoma-directed immunotherapy.
AU McLaughlin, Pamela M. J.; Harmsen, Martin C.; Dokter, Wim H. A.; Kroesen, Bart-Jan; van der Molen, Henk; Brinker, Marja G. L.; Hollerna, Harry; Ruiters, Marcel H. J.; Buys, Charles H. C. M.; de Leij, Lou F. M. H.
 CS Department of Pathology and Laboratory Medicine, Section Medical Biology,
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APPLICATION NO.

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Neutrania

I.f.m.h.de.leij@med.rug.nl

SO Cancer Research, (May 15, 2001) Vol. 61, No. 10, pp. 4105-4111. print.

CODEN: CNREA8. ISSN: 0008-5472.
DT Article
 LA English
ED Entered STN: 20 Jun 2001
       Last Updated on STN: 19 Feb 2002
Last Updated on S1N: 19 Feb 2002

AB The human pancarcinoma-associated ***epithelial***

***glycoprotein*** - ***2*** (EGP-2), a Mr 38,000 transmembrane
antigen also known as ***17*** - ***1A*** or ***Ep***

****CAM***, is commonly used for targeted immunotherapy of carcinomas
because it is strongly expressed by most carcinomas. EGP-2 is, however,
       also expressed in most normal epithelia. To evaluate anti-EGP-2-directed treatment-associated effects on tumors and on EGP-2-positive normal
       tissue, we generated EGP-2-expressing transgenic mice. A 55-kb DNA
       fragment consisting of the 14-kb genomic coding sequence of the human EGP-2 gene with apprx10-kb-upstream and apprx31-kb-downstream sequences
       was isolated and used to direct EGP-2 expression in an epithelium-specific manner. In the EGP-2 transgenic mice, EGP-2 appeared to be specifically expressed in all of those epithelial tissues that also express EGP-2 in
       humans, whereas all of the other tissues were negative. The specific in vivo localization of the i.v. administered anti-EGP-2 monoclonal antibody
       MOC31 was studied in EGP-2-positive and -negative tumors induced s.c. in
       this EGP-2 transgenic mouse model. Immunohistochemical analysis showed specific localization of MOC31 in the EGP-2-positive tumors but not in the
       SGP-2-negative tumors. No anti-EGP-2 monoclonal antibody localization was observed in any of the EGP-2-positive normal mouse tissues, which indicated a limited in vivo accessibility. In conclusion, an EGP-2
       transgenic mouse model has been generated that expresses the EGP-2 antigen as in humans and, therefore, can serve as a model to evaluate the efficacy and safety of a variety of anti-EGP-2-based immunotherapeutic modalities
       in both tumors and normal tissue
 L3 ANSWER 20 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
 AN 2000:881321 CAPLUS
 DN 134:38630
 TI Streptavidin expressed gene fusions forming tetrameric complexes with
therapeutic implications for adenocarcinomas and hematol. malignancies IN Goshorn, Stephen Charles; Graves, Scott Stoll; Schultz, Joanne Elaine;
       Lin, Yukang; Sanderson, James Allen; Reno, John M.
PA Neorx Corp., USA
SO PCT Int. Appl., 99 pp.
CODEN: PIXXD2
DT Patent
   A English
FAN.CNT 5
       PATENT NO.
                                                  KIND DATE
                                                                                          APPLICATION NO.
                                                                                                                                                DATE
PI WO 2000075333
      WO 2000075333
A1 20001214 WO 2000-US15595 20000605
WO 2000075333
C2 20020620
W. AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW. GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
CA 2376192 AA 20001214 CA 2000-2376192 20000605
EP 1190061 A1 20020327 EP 2000-941246 20000605
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
                                                                    20001214 WO 2000-US15595
                                                                                                                                                      20000605
                                                         A1
                   IE, SI, LT, LV, FI, RO
PRAI US 1999-168976P P 19991203
WO 2000-US15595 W 20000605
                                                                                                                                           20000605
 AB The present invention provides vectors for expressing genomic streptavidin
      fusion cassettes which include inducible promoters and various linkers and signal sequences. In the various embodiments, fusion proteins produced
       from these vectors are provided. In particular embodiments, fusion proteins comprising a single chain antibody (huNR-LU-10) and genomic streptavidin are provided as are vectors encoding the same. Also
       provided, are methods of using the fusion proteins of the present invention, and in particular, the use of scFvSA fusion proteins involving B9E9 as diagnostic markers or as a cell specific targeting agents. In
       addn. tetravalent antibodies that contact a fusion protin forming a tetrametric complex which may comprise a tumor cell surface-assocd.
      protein and a streptavidin portion capable of binding biotin and a biotinylated radionuclide contg. compd. A immunoreactivity assay is described in addn. to monitoring of blood clearance and turnor uptake of
      fusion proteins. Some adenocarcinomas and hematol, malignancies such as non-Hodgkin's lymphoma may be treated with these fusio-protein expressing
      non-hodgkin's symphoma may be treated with these fusio-protein expressin vectors. This system offers the expression of a genomic streptavidin gene fusion as a sol. protein into the periplasmic space of Escherichia coli where it undergoes spontaneous folding. This expression offers efficient protein folding where one does not need to purify and refold the protein
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THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD

University Hospital Groningen, Hanzeplein 1, 9713 GZ, Groningen,

Netherlands

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DN 134:16539
TI Antibodies
 IN Hoogenboom, Hendricus Renerus Jacobus Mattheus; Reurs, Anneke; Beiboer,
      Sigrid Herma Wilma
Oxford Biomedica (UK) Limited, UK
SO PCT Int. Appl., 86 pp.
CODEN: PIXXD2
DT Patent
 LA English
FAN.CNT 1
        PATENT NO.
                                                  KIND DATE
                                                                                          APPLICATION NO.
                                                                                                                                                 DATE
                                                                  20001123 WO 2000-GB1910
PI WO 2000069914
                                                                                                                                                      20000518
           /O 2000069914 A3 20010405
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
       WO 2000069914
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRAI GB 1999-11569 A 19990518

B Human antibodies that recognize the epithelial olycoprotein antigen
AB Human antibodies that recognize the epithelial glycoprotein antigen (EGP-2) are disclosed. The antibodies have a human light chain variable
        region and a human heavy chain variable region. Fragments of the
       antibodies and pharmaceutical compns. comprising the antibodies and their in vitro and in vivo applications in diagnosis and immunotherapy are also
         ANSWER 22 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
AN 2000:402017 CAPLUS.
DN 133:54574
        Recombinant vectors expressing multiple costimulatory molecules, host cell
       infection, and uses in immunogenic applications
Schlom, Jeffrey, Hodge, James, Panicali, Dennis
          United States Dept. of Health and Human Services, USA; Therion Biologics
       Corporation
PCT Int. Appl., 188 pp.
      CODEN: PIXXD2
DT Patent
  _A English
FAN.CNT 1
        PATENT NO.
                                                  KIND DATE
                                                                                          APPLICATION NO.
           WO 2000034494 A1 20000615 WO 1999-US26866 19991112
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
       WO 2000034494
          IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

A 2354024 AA 20000615 CA 1999-2354024 19991112

P 1137792 A1 20011004 EP 1999-958951 19991112

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

P 2002531133 T2 20020924 JP 2000-586927 19991112
       CA 2354024
EP 1137792
IE, SI, LI, LV, FI, RO
JP 2002531133 T2 20020924 JP 2000-586927
AU 774076 B2 20040617 AU 2000-16218
US 2004019195 A1 20040129 US 2003-406317
PRAI US 1998-111582P P 19981209
WO 1999-US26866 W 19991112
                                                                                                                                         19991112
                                                                                                                                               20030404
                                                     A3 20010924
        US 2001-856988
      3 The present invention provides recombinant vectors encoding and expressing at least three or more costimulatory mols and host cells infected by the vector. The recombinant vector may addnl. contain a gene encoding one or
        more target antigens or immunol. epitope as well as cytokine, chemokine, 
or Fit-3L. A method of making a recombinant poxvirus, of enhancing an 
immune response of an individual by administering a recombinant vector,
      and of treating or preventing a disease by activating a T lymphocyte, are also presented. Further describes are a method of making a progenitor
       dendritic cell or dendritic cell, of assessing the efficacy of a vaccine against a target antigen, and of screening for novel immunogenic peptides. The synergistic effect of these costimulatory mols. on the enhanced
       activation of T cells was demonstrated. The degree of T-cell activation using recombinant vectors contg. genes encoding three costimulatory mols. was far greater than the sum of recombinant vector constructs contg. one
       costimulatory mol. and greater than the use of two costimulatory mols. Results employing the triple costimulatory vectors were most dramatic
        under conditions of either low levels of first signal or low stimulator to 
T-cell ratios. This phenomenon was obsd. with both isolated CD4+ and CD8+
        T cells. The recombinant vectors of the present invention are useful as
      immunogenes and vaccines against cancer and pathogenic micro-organisi 
and in providing host cells, including dendritic cells and splenocytes 
with enhanced antigen-presenting functions.

E.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
RE.CNT 4
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L3 ANSWER 21 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN

2000;824304 CAPLUS

#### ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L3 ANSWER 23 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
  AN 2000:240985 CAPLUS
 DN 132:292701
  TI Novel methods for therapeutic vaccination
 IN Steinaa, Lucilla; Mouritsen, Soren; Nielsen, Klaus Gregorious; Haaning
       Jesper, Leach, Dana; Dalum, Iben; Gautam, Anand; Birk, Peter, Karlsson,
 PA M & E Biotech A/S, Den. SO PCT Int. Appl., 220 pp.
      CODEN: PIXXD2
 DT Patent
  LA English
 FAN.CNT 1
      PATENT NO.
                                                                APPLICATION NO.
                                    KIND DATE
                                                                                                      DATE
         WO 2000020027 A2 20000413 WO 1999-DK525 19991005
WO 2000020027 A3 20001012
W: AE, AL, AM, AT, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, CZ, DE, DE, DK, DK, DM, EE, EE, ES, FI, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK, LT, JT, MT, RT, TT, UA, UG, US, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
A2345817 AA 20000413 CA 1999-2345817 19991005
U 751709 B2 20020822
                                               20000413 WO 1999-DK525
 PI WO 2000020027
                                         A2
                                                                                                         19991005
      WO 2000020027
      CA 2345817
      AU 9958510
                                  B2
A2
                                          20020822
20010725
      AU 751709
                                                               EP 1999-945967
                                                                                                  19991005
      EP 1117421
      EP 1117421
                                    B1
                                           20040616
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, MC, IE, SI, LT, LV, FI, RO
      TR 200100936
                                             20010821 TR 2001-200100936
                                                                                                      19991005
      JP 2002526419
                                      T2
                                             20020820 JP 2000-573386
                                                                                                   19991005
      EE 200100203
                                             20021015 EE 2001-203
                                                                                                19991005
                                        20031031 NZ 1999-511055
20040715 AT 1999-945967
      NZ 511055
                                                                                                19991005
                                  E
      AT 269100
                                                                                                19991005
      NO 2001001586
                                              20010531 NO 2001-1586
                                                                                                   20010328
      ZA 2001002603
HR 2001000319
                                             20020930 ZA 2001-2603
20020630 HR 2001-319
                                                                                                 20010329
                                                                                                  20010504
                                              20040722 US 2003-441779
19981005
      US 2004141958
                                                                                                     20030519
                                       A
 PRAI DK 1998-1261
      US 1998-105011P
                                               19981020
                                       Å1
W
      US 1999-413186
                                               19991005
      WO 1999-DK525
                                                19991005
  AB A method is disclosed for inducing cell-mediated immunity against cellular
      antigens. More specifically, the invention provides for a method for inducing cytotoxic T-lymphocyte immunity against weak antigens, notably
      self-proteins. The method entails that antigen presenting cells are induced to present at least one CTL epitope of the weak antigen and at the same time presenting at least one foreign T-helper lymphocyte epitope. In
      a preferred embodiment, the antigen is a cancer specific antigen, e.g. prostate specific membrane antigen (PSM), Her2, or FGF8b. The method can
      be exercised by using traditional polypeptide vaccination, but also by
      using live attenuated vaccines or nucleic acid vaccination. The invention furthermore provides immunogenic analogs of PSM, Her2 and FGF8b, as well
     as nucleic acid mols. encoding these analogs. Also vectors and transformed cells are disclosed. The invention also provides for a method
      for identification of immunogenic analogs of weak or non-immunogenic
 L3 ANSWER 24 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 1999:795994 CAPLUS
 TI Gene probes used for genetic profiling in healthcare screening and
      planning
      Roberts, Gareth Wyn
 PA Genostic Pharma Ltd., UK
SO PCT Int. Appl., 745 pp.
     CODEN: PIXXD2
DT Patent
 LA English
 FAN.CNT 2
     PATENT NO.
                                    KIND DATE
                                                               APPLICATION NO.
                                                                                                      DATE
         VO 9964627 A2 19991216 WO 1999-GB1780 19990604
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             7. A.E., ALI, AM, AI, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ
DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRAI GB 1998-12099 A 19980606
      GB 1998-13291
                                             19980620
     GB 1998-13611
GB 1998-13835
                                            19980624
19980627
     GB 1998-14110
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GB 1998-15574
                                             19980718
    GB 1998-15576
GB 1998-16085
                                             19980718
                                             19980724
                                     AAAA
      GB 1998-16086
                                             19980724
      GB 1998-16921
                                             19980805
     GB 1998-17097
                                             19980807
     GB 1998-17200
                                             19980808
      GB 1998-17632
                                             19980814
      GB 1998-17943
                                            19980819
AB There is considerable evidence that significant factor underlying the
     individual variability in response to disease, therapy and prognosis lies
      in a person's genetic make-up. There have been numerous examples relating
    that polymorphisms within a given gene can alter the functionality of the protein encoded by that gene thus leading to a variable physiol, response
    In order to bring about the integration of genomics into medical practice and enable design and building of a technol. platform which will enable the everyday practice of mol. medicine a way must be invented for the DNA sequence data to be aligned with the identification of genes central to the induction, development, progression and outcome of disease or physiol. states of interest. According to the invention, the no. of genes and
    their configurations (mutations and polymorphisms) needed to be identified in order to provide crit. clin. information concerning individual
     prognosis is considerably less than the 100,000 thought to comprise the
    human genome. The identification of the identity of the core group of genes enables the invention of a design for genetic profiling technologies
    genes enables the invention of a design for geneue profiling technologies which comprises of the identification of the core group of genes and their sequence variants required to provide a broad base of clin. prognostic information - "genostics". The "Genostic" profiling of patients and persons will radically enhance the ability of clinicians, healthcare professionals and other parties to plan and manage healthcare provision
     and the targeting of appropriate healthcare resources to those deemed most
    in need. The use of this invention could also lead to a host of new applications for such profiling technologies, such as identification of
     persons with particular work or environment related risk, selection of
    applicants for employment, training or specific opportunities or for the enhancing of the planning and organization of health services, education
     services and social services.
L3 ANSWER 25 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
AN 1999:795993 CAPLUS
      132:31743
TI Gene probes used for genetic profiling in healthcare screening and
    planning
Roberts, Gareth Wyn
PA Genostic Pharma Limited, UK
SO PCT Int. Appl., 149 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 2
                                  KIND DATE
                                                              APPLICATION NO.
                                                                                                    DATE
    PATENT NO.
    WO 9964626
        VO 9964626 A2 19991216 WO 1999-GB1779 19990604
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
            DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ
            TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ. TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     CA 2330929
                                          19991216 CA 1999-2330929
                                                                                                  19990604
    AU 9941586
                                         19991230 AU 1999-41586
                                  A1
                                                                                                19990604
     AU 766544
                                         20031016
    AU 9941587
GB 2339200
                                  Α1
                                          19991230
20000119
                                                             AU 1999-41587
                                                                                                19990604
                                  A1
                                                             GB 1999-12914
                                                                                                19990604
     GB 2339200
                                          20010912
    EP 1084273 A1 20010321 EP 1999-925207 19990604
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
    IE, FI
JP 2003528564
                                           20030930 JP 2000-553616
20031023 US 2002-206568
                                                                                                  19990604
     US 2003198970
                                     A1
                                                                                                    20020729
PRAI GB 1998-12098
GB 1998-28289
                                        Α
                                               19980606
                                           19981223
    GB 1998-16086
GB 1998-16921
                                           19980724
19980805
    GB 1998-17097
                                            19980807
    GB 1998-17200
GB 1998-17632
                                           19980808
19980814
     GB 1998-17943
                                           19980819
     US 1999-325123
                                     B1
                                             19990603
    WO 1999-GB1779
                                       W
                                              19990604
AB There is considerable evidence that significant factor underlying the
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individual variability in response to disease, therapy and prognosis lies in a person's genetic make-up. There have been numerous examples relating that polymorphisms within a given gene can alter the functionality of the protein encoded by that gene thus leading to a variable physiol. response. In order to bring about the integration of genomics into medical practice

and enable design and building of a technol, platform which will enable the everyday practice of mol. medicine a way must be invented for the DNA

GB 1998-14580 GB 1998-15438 19980707 19980716

sequence data to be aligned with the identification of genes central to the induction, development, progression and outcome of disease or physiol. states of interest. According to the invention, the no. of genes and their configurations (mutations and polymorphisms) needed to be identified in order to provide crit. clin. Information concerning individual prognosis is considerably less than the 100,000 thought to comprise the human genome. The identification of the identity of the core group of genes enables the invention of a design for genetic profiling technologies.

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L3 ANSWER 26 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
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1999:691109 CAPLUS

DN 131:335805

TI Glycosylation engineering of antibodies for improving antibody-dependent cellular cytotoxicity

Umana, Pablo; Jean-Mairet, Joel; Bailey, James E.

Switz.

SO PCT Int. Appl., 79 pp. CODEN: PIXXD2

DT Patent

LA English FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

A1 19991028 WO 1999-US8711 PI WO 9954342 19990420 

TM, TR, TT, UA, UG, US, UZ, VN, TO, Z-1, ..., MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

U 9936578 A1 19991108 AU 1999-36578 19990420

P 1071700 A1 20010131 EP 1999-918731 19990420

TO CALL DE DY ES FR GB GR, IT, LI, LU, NL, SE, MC, PT, AU 9936578 EP 1071700 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE ÉL

T2 20020423 JP 2000-544680 B1 20030805 US 1999-294584 A1 20040415 US 2003-437388 P P 19980420 JP 2002512014 19990420 US 6602684 19990420 US 2004072290 20030514

PRAI US 1998-82581P US 1999-294584 A1 19990420 W 19990420 WO 1999-US8711

AB The present invention relates to the field of glycosylation engineering of proteins. More particularly, the present invention is directed to the glycosylation engineering of proteins to provide proteins with improve therapeutic properties, e.g., antibodies, antibody fragments, or a fusion protein that includes a region equiv. to the Fc region of an Ig, with enhanced Fc-mediated cellular cytotoxicity. The antibodies or fusion proteins with enhanced Fc-mediated cellular cytotoxicity are expressed in host cells engineered to also express a glycoprotein-modifying glycosyl transferase, e.g. .beta.(1,4)-N-acetylglucosaminyltransferase III or V, .beta.(1,4)-N-galactosyltransferase, and mannosidase II. .CNT 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS

RECORD

### ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 27 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN

AN 1998:173463 CAPLUS

DN 128:304704

TI A -308 deletion of the tomato LAP promoters is able to direct flower-specific and MeJA-induced expression in transgenic plants

AU Ruiz-Rivero, Omar J.; Prat, Salome

CS Dpto. de Genetica Molecular, Centro de Investigacion y Desarrollo-C.S.I.

C., Barcelona, 08034, Spain SO Plant Molecular Biology (1998), 36(5), 639-648 CODEN: PMBIDB; ISSN: 0167-4412

PB Kluwer Academic Publishers

DT Journal

AB Tomato and potato leucine aminopeptidase (LAP) mRNAs are induced in response to mech. wounding and the wound signal mols., ABA and jasmonic acid. Here, we report the isolation of two LAP genes, LAP17.1A and LAP17.2, from tomato. Functional anal. in transgenic tomato and potato plants show that fusions of the corresponding 5' non-coding regions to the gusA gene are constitutively expressed in flowers and induced in leaves upon wounding or by treatment with Me jasmonate (MeJA). Comparison of the upon wounding regions of the two genes revealed a region from -317 to -3 relative to the ATG, which is strongly conserved in both promoters. This 0.3 kb proximal \*\*\*promoter\*\*\* fragment is sufficient to direct flower-specific and MeJA-inducible GUS activity in transgenic potato plants, and thus contains a MeJA-responsive element that mediates induction by MeJA. Dimeric TGACG motifs or G-box elements similar to those found in other MeJA-inducible genes are not obsd. in this region, which suggests that a different DNA sequence is involved in MeJA induction

of the LAP genes.

RE.CNT 37 THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 28 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation, on

**DUPLICATE 4** 

1998:436540 BIOSIS DN PREV199800436540

The impact of antigen density and antibody affinity on antibody-dependent

cellular cytotoxicity: Relevance for immunotherapy of carcinomas.

AU Velders, M. P.; Van Rhijn, C. M.; Oskam, E.; Fleuren, G. J.; Warnaar, S. O.; Litvinov, S. V. [Reprint author]

CS Dep. Pathol., Leiden Univ. Hosp. Build. 1, L1-Q, PO Box 9600, 2300 RC

SO British Journal of Cancer, (Aug., 1998) Vol. 78, No. 4, pp. 478-483.

CODEN: BJCAAI. ISSN: 0007-0920.

DT Article LA English

ED Entered STN: 7 Oct 1998

Last Updated on STN: 7 Oct 1998

AB Antibody-dependent cellular cytotoxicity (ADCC) is considered to be the major mechanism through which tumour cells, upon treatment with anti-tumour MAbs, are eliminated in vivo. However, the relative importance of various parameters that influence the efficacy of ADCC is unclear. Here we present in vitro data on the impact of MAb affinity and unclear. Here we present in vitor data on the impact of man anning and antigen density on ADCC, as obtained by comparison of two MAbs against the tumour-associated antigen \*\*\*Ep\*\*\* - \*\*\*CAM\*\*\* . The low-affinity MAb \*\*\*17\*\*\* - \*\*\*1A\*\*\* (Ka = 5 X 107 M-1) currently used for therapy, and the high-affinity MAb 323/A3 (Ka = 2 X 109 m-1), were compared in ADCC experiments against murine and human turnour target cells transfected with the \*\*\*Ep\*\*\* - \*\*\*CAM\*\*\*\* cDNA under the control of an inducible \*\*\*promoter\*\*\* to enable regulation of the target antigen an induction promoter to enable regulation of the larget analyse expression levels. Data obtained from these studies revealed that the high-affinity MAb, in contrast to the low-affinity MAb, could mediate killing of turnour cells with low antigen expression levels. Even at comparable MAb-binding levels, ADCC mediated by the high-affinity MAb was more effective. The kinetics of ADCC was also found to be determined by the level of antigen expression, and by the affinity and the concentration of the MAb used. The efficacy of ADCC with both low- and high-affinity MAbs further depended on adhesive interactions between effector and target cells mediated by CD18. However, at every given MAb concentration these interactions were of less importance for the high-affinity MAb than for the low-affinity MAb. As heterogeneity of a target antigen expression is a common feature of all tumours, and some tumour cells express very low levels of the antigen, the use of high-affinity MAbs in immunotherapy may significantly improve the clinical results obtained to the present date in the treatment of minimal residual disease.

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L3 ANSWER 29 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 1997:97727 CAPLUS
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126:156420

TI Prophylactic and therapeutic vector vaccination using expression constructs for individual epitopes of antigens

IN Weiner, David B.; Williams, William V.; Wang, Bin
PA Wistar Institute, USA; Trustees of the University of Pennsylvania
SO U.S., 50 pp., Cont.-in-part of U.S. Ser. No. 29,336, abandoned.
CODEN: USXXAM

DT Patent LA English FAN.CNT 4

> PATENT NO. KIND DATE APPLICATION NO. DATE US 5593972 19970114 US 1993-125012 19930921 19950103 ZA 1994-493 19940804 CA 1994-2153593 19940804 WO 1994-US899 19940125 ZA 9400493 19940126 CA 2153593 WO 9416737 19940126 /O 9416737 A1 19940804 WO 1994-US899 19940126
> W: AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU, JP, KP, KR, KZ, LK, LU, LV, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SK, UA, US, US, US, US, US, UZ, VN
> RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
> U 9462320 A1 19940815 AU 1994-62320 19940126
> U 675702 B2 19970213 AU 9462320 AU 675702

EP 681483 A1 19951115 EP 1994-909492 19940126 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE HU 73099 A2 19960628 HU 1995-2229 19940126 HU 219767 20010730 T2 A2 JP 08509694 19961015 JP 1994-517285 19940126 EP 1473369 20041103 EP 2004-75092 19940126 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL US 6348449 B1 20020219 US 1994-357398 SE, MC, PT, IE 19941216 A A B1 US 1995-453349 US 1997-783818 US 5830876 19981103 US 5817637 US 6468982 19981006 19970113 20021022 US 1997-880576 19991109 US 1997-979385 B2 19930126 19970623

19971126

US 5981505 PRALUS 1993-8342 US 1993-29336 B2 19930311 US 1993-93235 US 1993-124962 19930715 19930921 US 1993-125012 EP 1994-909492 19930921 A3 19940126

WO 1994-US899 19940126 US 1995-495684 В1 19950828 US 1997-783818 19970113 A1

AB Methods of prophylactic and therapeutic immunization against infection, hyperproliferative and autoimmune diseases are disclosed. An expression construct directing the synthesis of one or more epitopes, or analogs of

epitopes, of an antigen is introduced into cells of an individual. The epitope is identical or substantially similar to an epitope of a pathogen antigen, a hyperproliferative cell assocd. protein or a protein assocd. with autoimmune disease resp. Methods of immunizing against HIV are described. Successful induction of immunity to HIV1 in mice by injection with an expression vector for the HIV-1 gene env.

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L3 ANSWER 30 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
AN 1996:278128 CAPLUS
DN 124:307777
TI Dynamic monitoring and quantification of gene expression in single, living
cells: a molecular basis for secretory cell heterogeneity
AU Castano, Justo P.; Kineman, Rhonda D.; Frawley, L. Stephen
CS Dep. Cell Biology Anat., Med. Univ. South Carolina, Charleston, SC, 29425,
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SO Molecular Endocrinology (1996), 10(5), 599-606 CODEN: MOENEN; ISSN: 0888-8809

PB Endocrine Society DT Journal

USA

LA English

AB Progress in understanding the dynamics of gene expression has been hampered by lack of a strategy for continuously monitoring this process within normal, living cells. Here, the authors employed a modifn. of conventional luciferase technol. to make single and repeated real-time measurements of PRL gene expression from individual, living lactotropes from nursing rats. Cells were individually transfected by microinjection with a PRL \*\*\*promoter\*\*\* /luciferase reporter construct. Levels of PRL gene transcription were quantified by photonic imaging in the same cells before and after 24 h of culture in the presence or absence of the dopamine agonist bromocryptine or \*\*\*EGF\*\*\*, \*\*\*2\*\*\* well known regulators of PRL gene transcription. These cells were found to be remarkably heterogeneous with respect to basal PRL gene expression and that the degree of activity within a single cell could fluctuate greatly over time under basal culture conditions. Treatment with bromocryptine or EGF induced predictable and reversible changes in the av. responses obsd., yet individual cells displayed marked differences in responses to these agents. These findings demonstrate the utility of this paradigm for monitoring dynamics of gene expression within normal, living cells of any type. Moreover, they provide a mol. basis for the secretory heterogeneity and plasticity that have come to be known as hallmarks of lactotrope cell

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L3 ANSWER 31 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN
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1995:319826 CAPLUS

DN 122:98808

T1 Cloning and expression of human .beta.2-microglobulin cDNA and the construction of fusion proteins between antigenic epitopes and

.beta.2-microglobulin IN Edwards, Richard Mark; Hunter, Michael George

PA British Bio-Technology Ltd., UK SO PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN CNT 2

PATENT NO. KIND DATE APPLICATION NO. DATE PI WO 9424290 

W. AU, BR, CA, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE AU 9464353 A1 19941108 AU 1994-64353 19940408 EP 693125 A1 19960124 EP 1994-91204 19940408 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, LU, NL, PT, SE US 200212318 A1 20020905 US 1995-532549 19951201

PRAI GB 1993-7371 19930408 w WO 1994-GB755 19940408

AB A method is described for the cloning and expression of human beta.2-microglobulin (B2M) cDNA in vector host cells which allows the construction of B2M fusion proteins with antigenic sequences from various etiol, agents or tumors. Preferred antigenic sequences are derived from the third variable domain (V3 loop) of an envelope protein of a lentivirus. These fusion proteins can be used as prophylactic or immunotherapeutic vaccines to induce neutralizing antibody response immunotherapeutic vaccines to induce neutralizing antibody responses. Thus, B2M cDNA was inserted into the pHILD1 expression vector for expression in the Pichia pastoris system. The expression vector includes an AOX \*\*\*promoter\*\*\* sequence and an .alpha.-factor or Pho1 leader sequence to obtain secretion of the fusion protein from the yeast cells. Within the Pichia pastoris expression system, the B2M gene was fused at its 5' end to the Sendai virus epitope (FAPGNYPAL-GGGGG, where the pentaglycine is a short linker) or to the influenza A virus nucleoprotein epitope (GILGFVFTL-GGGGGGSSS). Prodn. levels from strains with the alpha.-factor leader sequence were .apprx.150 mg/L. The hybrid Sendai-B2M product was shown to induce Sendai nucleoprotein-specific cytotoxic T-lymphocytes.

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L3 ANSWER 32 OF 37 CAPLUS COPYRIGHT 2005 ACS on STN AN 1994:623662 CAPLUS
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DN 121:223662

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SO PCT Int. Appl., 135 pp.
    CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 4
    PATENT NO.
                                KIND DATE
                                                          APPLICATION NO.
                                                                                              DATE
PI WO 9416737
                                  A1 19940804 WO 1994-US899
                                                                                              19940126
        W: AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU, JP, KP, KR, KZ, LK, LU, LV, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SK, UA, US, US, US, US, US, UZ, VN
     RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
US 5593972 A 19970114 US 1993-125012 19930921
ZA 9400493 A 19950103 ZA 1994-493 19940125
CA 2153593 AA 19940804 CA 1994-2153593 19940126
    ZA 9400493
     AU 9462320
AU 675702
                                      19940815 AU 1994-62320
19970213
                                                                                         19940126
                               B2
                                                                                         19940126
                                       19951115 EP 1994-909492
     EP 681483
     R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
JP 08509694 T2 19961015 JP 1994-517285 19940126
RU 2174845 C2 20011020 RU 1995-117922 19940126
     RU 2174845
US 5981505
                                      19991109 US 1997-979385
                                Α
                                                                                         19971126
PRAI US 1993-8342
                                           19930126
     US 1993-29336
                                  Α
                                         19930311
    US 1993-93235
US 1993-124962
                                        19930715
    US 1993-125012
WO 1994-US899
                                   A
W
                                         19930921
19940126
     US 1995-495684
                                          19950828
AB Methods of introducing nucleic acids into cells of an individual using
     agents that stimulate nucleic acid uptake or expression or the
     inflammatory response are described. The method avoids the use of viral
    or retroviral particles. The transforming nucleic acid encodes an antigenic peptide and so may be useful in therapeutics or prophylaxis
    Methods of prophylactically and therapeutically immunizing an individual against HIV without the use of retroviral proteins or particles are
     disclosed. Expression cassettes for manuf. of antigens of HIV-1 in animal
    cells were constructed by std. methods. These were used to transform tumor cell lines not normally recognized by a mouse host. Mice injected
     with these transformed cells mounted a strong cytotoxic response that
    completely eliminated tumors that would normally kill the animal in 12 wk. Injection of mice with an expression vector carrying an expression
     cassette for gp160 in combination with bupivacaine to stimulate
    inflammation and cell proliferation resulted in a strong immune response to gp160. The response was stronger than from mice injected with gp160 or
    injected with the expression vector without the use of bipuvacaine
L3 ANSWER 33 OF 37 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.
                                                        DUPLICATE 5
AN 95036827 EMBASE
DN 1995036827
TI Studies on **17*** - ***1A**** antigen gene regulation in nonexpressing 549 and A431 cells, as compared to expressing pancreatic
     carcinoma (Capan 2) cells, reveal a complex mechanism of repression of
AU Siemieniako B.; Wiland E.; Trzeciak W.H.
CS Inst. of Biochemistry/Biotechnology, University of Agriculture, Wolynska
    35,60-637 Poznan, Poland
SO Cell Biology International, (1994) 18/11 (1009-1017).
ISSN: 1065-6995 CODEN: CBIIEV
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CY United Kingdom DT Journal; Article

FS 016 Cancer 022 Human Genetics 029 Clinical Biochemistry

English

SL English

Elements controlling high expression of the \*\*\*17\*\*\* - \*\*\*1A\*\*\* antigen gene in pancreatic carcinoma cells (Capan 2) reside within the two regions: proximal (-193 to +3) and distal (-877 to -518). We demonstrate here that some factors present in nuclear extracts from nonexpressing cells bind specifically to the control elements, important for gene expression. Our results suggest that nonexpressing cells may either lack at least one of the factors necessary for activation or may contain their modified forms. A major difference between expressing and nonexpressing recells was found in the region containing core enhancer sequence. Moreover, nonexpressing cells display a complex pattern of DNA-protein interactions in this region, suggesting that these cells contain factors acting negatively mainly on the enhancer sequence. Our results however, indicate that the mechanism of repression is much more complicated than expected.

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L3 ANSWER 34 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
Corporation. on
                              DUPLICATE 6
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AN 1993:207235 BIOSIS

DN PREV199395108460

TI Retroposition in a family of carcinoma-associated antigen genes.
AU Linnenbach, Alban J. (Reprint author); Seng, Beth A.; Wu, Shuang; Robbins,
Shira; Scollon, Maureen; Pyrc, Jania J.; Druck, Teresa; Huebner, Kay
CS Wistar Inst., 3601 Spruce St., Philadelphia, PA 19104, USA
SO Molecular and Cellular Biology, (1993) Vol. 13, No. 3, pp. 1507-1515.

TI Genetic transformation of animal cells using agents that stimulate DNA uptake or gene expression or the inflammatory response IN Weiner, David B.; Williams, William V.; Wang, Bin; Coney, Leslie R.; Merva, Michael J.; Zurawski, Vincent R., Jr.

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CODEN: MCEBD4. ISSN: 0270-7306.
 DT Article
LA English

    OS Genbank-M93029; Genbank-M93030; Genbank-M93031; Genbank-M93032;
    Genbank-M93033; Genbank-M93034; Genbank-M93035; Genbank-M93036;

       Genbank-X13425
 ED Entered STN: 23 Apr 1993
Last Updated on STN: 9 Jun 1993
  AB The gene encoding the carcinoma-associated antigen defined by the
     monoclonal antibody GA733 is a member of a family of at least two type I membrane proteins. This study describes the mechanism of evolution of the GA733-1 and ****GA733*** - ********* genes. A full-length cDNA clone for GA733-1 was obtained by screening a human placental library with a
      genomic DNA probe. Comparative analysis of the cDNA sequence with the
      previously determined genomic sequence confirmed that GA733-1 is an intronless gene. The ***GA733*** - ***2*** gene encoding the
     intronless gene. The ""GA733" - ""2" gene encoding the monoclonal antibody-defined antigen was molecularly cloned with a cDNA probe and partially sequenced. Comparison of ""GA733" - ""2" gene sequences with the previously established cDNA sequence revealed that this gene consists of nine exons. The putative ""promoter" regions of the GA733-1 and ""GA733" - ""2"" genes are unrelated. These findings suggest that the GA733-1 gene was formed by the retroposition of the ""GA733" - ""2" gene via an mRNA intermediate. Prior to retroposition, the ""GA733" gene had been affected by exon shuffling. Analysis of ""GA733" gene had been affected by exon shuffling. Analysis of ""GA733". The GA733-1 retroposon was localized either to chromosome region 1p32-1p31 or to 1p13-1q12, and the ""GA733" - ""2" founder gene was localized to chromosome 4q.
 L3 ANSWER 35 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
 Corporation. on
                                                            DUPLICATE 7
     STN
 AN 1992:476438 BIOSIS
DN PREV199294107813; BA94:107813
  TI NUCLEAR PROTEINS FROM CAPAN-2 CELL LINE FORM SPECIFIC
 COMPLEXES WITH THE
17-1 A ANTIGEN GENE ***PROMOTER***
 AU SIEMIENIAKO B [Reprint author]; WILAND E
CS INST HUMAN GENETICS, POLISH ACADEMY SCI, STRZESZYNSKA 32,
     POLAND
 SO Biochemical and Biophysical Research Communications, (1992) Vol. 186, No.
      3, pp. 1353-1361.
      CODEN: BBRCA9, ISSN: 0006-291X.
 DT Article
FS BA
LA ENGLISH
 ED Entered STN: 27 Oct 1992
     Last Updated on STN: 27 Oct 1992
 AB To determine the location of sites important for the function of the ***17*** - ***1A*** antigen gene ***promoter*** and to
      characterize the protein factors binding to these sites, fragments of the
***promoter*** region were analysed by gel retardation assay with
      nuclear extracts from Capan 2 cell line. At least two separate regions,
      which specifically bind nuclear proteins were identified within the 5'flanking region of the ***17*** - ***1A*** antigen gene. These regions have been located between nucleotides -877 to -518 (distal region)
      and -193 to +3 (proximal region) and presumably participate in regulation of expression of the ***17*** - ***1A*** antigen gene.
 L3 ANSWER 36 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
Corporation. on STN
       1992:134089 BIOSIS
DN PREV199242061789; BR42:61789
TI HUMAN ***17*** - ***1A*** NEOANTIGEN GENE ***PROMOTER*** .
AU WOUCIEROWSKI J (Reprint author); POLUHA D; ZIELEWICZ J
CS DEP MED GENETICS, MED SCH, 20090-LUBLIN, 8 JACZEWSKI STR,
SO American Journal of Human Genetics, (1991) Vol. 49, No. 4 SUPPL, pp. 434.

Meeting Info.: PROCEEDINGS OF THE 8TH INTERNATIONAL CONGRESS
 OF HUMAN
     GENETICS, WASHINGTON, D.C., USA, OCTOBER 6-11, 1991. AM J HUM
     CODEN: AJHGAG. ISSN: 0002-9297.
DT Conference; (Meeting)
FS BR
LA ENGLISH
ED Entered STN: 5 Mar 1992
     Last Updated on STN: 5 Mar 1992
L3 ANSWER 37 OF 37 BIOSIS COPYRIGHT (c) 2005 The Thomson
Corporation. on STN
                                                           DUPLICATE 8
AN 1988:311405 BIOSIS
DN PREV198886028443; BA86:28443
      TRANSFORMING GROWTH FACTOR BETA AS A POTENT
  ***PROMOTER*** IN
     TWO-STAGE BALB-C 3T3 CELL TRANSFORMATION.
       HAMEL E [Reprint author]; KATOH F; MUELLER G; BIRCHMEIER W;
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CS INTERNATIONAL AGENCY RES CANCER, 150 COURS ALBERT THOMAS,

69372 LYON CEDEX

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SO Cancer Research, (1988) Vol. 48, No. 10, pp. 2832-2836.
     CODEN: CNREA8. ISSN: 0008-5472.
 DT Article
 FS BA
 LA ENGLISH
ED Entered STN: 3 Jul 1988
     Last Updated on STN: 3 Jul 1988
 AB We have tested transforming growth factor .beta. (TGF,beta.) in the
     two-stage BALB/c 3T3 cell transformation assay for possible tumor-promoting activity, since it has several effects similar to those of
     tumor-promoting phorbol ester. After initiation of BALB/c 3T3 cells with 3-methylcholanthrene, treatment with TGF.beta. at 1 ng/ml alone or in
    combination with epidermal growth factor (EGF) for 4 weeks enhanced the number of transformed foci by 5- to 6-fold in comparison with uninitiated cells. Initiation treatment alone induced no or very few transformed foci
     in several assays. Treatment with phorbol-12,13-didecanoate (PDD) at 100
     ng/ml for 4 weeks enhanced the number of transformed foci in initiated
     BALB/c 3T3 cells by 4- to 5-fold in comparison with uninitiated cells.
    Thus, TGF beta. at 1 ng/ml is as potent as PDD at 100 ng/ml for tumor-promoting activity in the two-stage BALB/c 3T3 cell transformation
     assay. The enhancing effect of TGF.beta. was dose-related inthe dose
     range tested (0.03-1 ng/ml) and was not reversible. Some of the foci induced by combined MCA-TGF.beta.-EGF treatment were cloned, and eight
    of nine clones tested produced tumors in nude mice. TGF.beta. (1 ng/ml) plus ***EGF*** ( ***2*** ng/ml) increased the saturation density to
     a similar extent as PDD (100 ng/ml) but did not affect the growth of
     BALB/c 3T3 cells. We observed no change in junctional intercellular
    communication, as measured by the dye transfer method, when cells were treated with TGF beta. during the two-stage BALB/c 3T3 cell transformation assay. Nevertheless, there was selective communication between
     transformed and surrounding nontransformed cells, MCA-TGF beta
    transformed cells intercommunicated among themselves but not with surrounding nontransformed cells. Our results indicate that TGF.beta. has
    potent tumor-promoting activity in vitro, but that this activity is not mediated by a complete blockage of intercellular communication, as is
     suggested for phorbol ester tumor promoters.
 => d his
     (FILE 'HOME' ENTERED AT 16:15:29 ON 19 JAN 2005)
    FILE 'BIOSIS, EMBASE, CAPLUS' ENTERED AT 16:15:40 ON 19 JAN 2005
           1526 S EGF 2 OR EPITHELIAL GLYCOPROTEIN 2 OR EP CAM OR 17
             52 S L1 AND (PROMOTER OR REGULA? ELEMENT OR REGULAT?
 REGION OR 5 U
            37 DUP REM L2 (15 DUPLICATES REMOVED)
=> s carcinoma (3a) (select? or restrict? or specific?)
1 FILES SEARCHED...
         5817 CARCINOMA (3A) (SELECT? OR RESTRICT? OR SPECIFIC?)
 => s I4 and (promoter or regula? element or regulat? region or 5 UTR)
          235 L4 AND (PROMOTER OR REGULA? ELEMENT OR REGULAT?
REGION OR 5
≈> s I5 and lung carcinoma
L6 6 L5 AND LUNG CARCINOMA
 => dup rem 16
 PROCESSING COMPLETED FOR L6
             4 DUP REM L6 (2 DUPLICATES REMOVED)
 YOU HAVE REQUESTED DATA FROM 4 ANSWERS - CONTINUE? Y/(N):y
 L7 ANSWER 1 OF 4 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation.
 DUPLICATE 1
AN 2001:225975 BIOSIS
 DN PREV200100225975
TI Adenovirus-mediated suicide gene transfer to small cell ***lung***

***carcinoma*** using a tumor- ***specific*** ***promoter***.
 AU Morimoto, Emiko; Inase, Naohiko [Reprint author]; Miyake, Shuji;
    Yoshizawa, Yasuvuki
CS Pulmonary Medicine, Tokyo Medical and Dental University, 1-5-45 Yushima,
Bunkyo-ku, Tokyo, 113-8519, Japan ninase pulm@tmd.ac.jp

SO Anticancer Research, (January-February, 2001) Vol. 21, No. 1A, pp. 329-331. print.

CODEN: ANTRD4. ISSN: 0250-7005.
DT Article
LA English
ED Entered STN: 9 May 2001
    Last Updated on STN: 18 Feb 2002

The gastrin-releasing peptide (GRP) is expressed in most types of small cell ***lung*** ***carcinoma*** (SCLC) and the GRP

***promoter*** is thought to be potentially useful for tumor-specific
    expression of the suicide gene in SCLC. We constructed an adenovirus
```

containing the herpes simplex thymidine kinase suicide gene driven by the GRP \*\*\*promoter\*\*\* (AdGRP-TK) and transfected it into GRP-expressing SCLC cells (SBC5) to confer sensitivity to gancidovir (GCV). After infection with AdGRP-TK, SBC5 cells became more sensitive to GCV in vitro. In nude mice, a subcutaneously-inoculated tumor of SBC5 cells infected with AdGRP-TK in advance regressed completely after intraperitoneal administration of GCV. These results suggest that adenovirus-mediated gene transfer of the suicide gene followed by pro-drug treatment may be applicable to SCLC.

L7 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN

AN 2001:294518 CAPLUS

DN 135:220767

TI Neuron specific enolase \*\*\*promoter\*\*\* for suicide gene therapy in small cell \*\*\*lung\*\*\* \*\*\*carcinoma\*\*\* \*\*\*lung\*\*\*

AU Tanaka, Michiko; Inase, Naohiko; Miyake, Shuji; Yoshizawa, Yasuyuki CS Pulmonary Medicine, Tokyo Medical and Dental University, Tokyo, 113-8519,

SO Anticancer Research (2001), 21(1A), 291-294 CODEN: ANTRD4; ISSN: 0250-7005

PB International Institute of Anticancer Research

DT Journal

AB To investigate the specific transduction of a suicide gene into human small cell \*\*\*\*lung\*\*\* \*\*\*carcinoma\*\*\*\* (SCLC) cells, we explored the \*\*\*\*promoter\*\*\* region of the neuron specific enclase (NSE) gene as a tumor-specific \*\*\*promoter\*\*\*. In Northern blot anal, NSE mRNA was expressed more abundantly in the SBC3 human SCLC cell line than in the RERF human SCLC cell line, the A549 human lung adenocarcinoma cell line and the HeLa human uterine cervix epitheloid carcinoma cell line. A reporting vector contg. the NSE \*\*\*promoter\*\*\* (pNSE-LUC) exhibited higher luciferase activity in SBC3 than in the other three cell lines.

After transfecting an expression vector contg. the NSE \*\*\*promoter\*\*\* After transfecting an expression vector contg. the NSE ""promoter""
-bound HSV-TK gene. (pNSE-TK) into the cells, we measured their
sensitivity to ganciclovir (GCV). In SBC3, pNSE-TK transfected cells
showed about the same sensitivity to GCV as non-transfected (parental)
cells. Though the NSE ""promoter"" itself is not optimal for use in
suicide gene transfer to SCLC cells, it might be applied as a
tumor-specific ""promoter"" after enhancement of its activity.

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN

AN 1999:119163 CAPLUS DN 131:3509

RECORD

- Specific point-mutate p53 mini-gene transfecting effects on biological behaviors of a human cancer cell line PG derived from human pulmonary giant carcinoma
- AU Xie, Jiarrwu; Fang, Weigang; Hui, Pei; Li, Baolin; Li, Hongmei; Zhong, Gaogao; Zheng, Jie; Chen, Bifen; Wu, Bingquan
  CS Department of Molecular and Biology, Fuzhou Medical University, Fuzhou,
- 350005, Peop. Rep. China SO Zhonghua Yixue Zazhi (1999), 79(1), 57-60 CODEN: CHHTAT; ISSN: 0376-2491

PB Zhonghua Yixue Zazhi

DT Journal

LA Chinese
AB The suppressive effects of a murine genomic p53 minigene contg. an Arg-Leu substitution at its encoding amino acid 172 on biol. behaviors of human carcinoma cell were explored and its potential application in cancer gene therapy was evaluated. This mutant p53 gene which lacked of exon 1 and intron 1 expression vector driven by CMV "\*\*rpormoter\*\* was co-transfected with PCMVneo into PG cell in which dominant neg. p53 pre-exists by LipofectaMINE and electroporation methods. A wild-type and another kind of genomic mutate-type p53 gene expression vector were transfected. The latter p53 gene encoding protein contained an Arg-His substitution at the same position, and pBLuscript plasmid was used as control. All transfectants were screened by 500 .mu.g/mL geneticin and identified by mouse specific p53 mRNA RT-PCR and Northern blot anal. The biol. behavior changes were studied by colony formation and TUNEL test together with in-situ clone regression for chemosensitivity of anti-cancer drugs after transfection. The transfecting effects of this unusual p53 gene were surprisingly strong. They were more significant than those of the wild-type p53 and could suppress the formation of transgenic colonies and passage. The transgenic colonies were sensitive to be treated in adriamycin and 5-Fu, and the gene transient expression could give cell apoptosis. Codon 172 mutant (Arg-Leu) p53 genomic DNA exhibited a strong suppressive transfecting effects on carcinoma cell, so it was a possible candidate to be used in cancer gene therapy.

L7 ANSWER 4 OF 4 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

AN 92191536 EMBASE

- DN 1992191536
  TI Identification of a negative \*\*\*regulatory\*\*\* \*\*\*element\*\*\* that inhibits c-mos transcription in somatic cells.
- AU Zinkel S.S.; Pal S.K.; Szeberenyi J.; Cooper G.M.
- CS Dana-Farber Cancer Institute, Harvard Medical School, Boston, MA 02115, **United States**
- SO Molecular and Cellular Biology, (1992) 12/5 (2029-2036). ISSN: 0270-7306 CODEN: MCEBD4

CY United States

Journal; Article 004 Microbiology FS

English

SI. English

AB We have used transient expression assays to identify a cis-acting region in the 5 flanking sequence of murine c-mos which, when deleted, allows expression from the c-mos \*\*\*\*promoter\*\*\* in NIH 3T3 cells. This negative regulatory sequence, located 400 to 500 nucleotides upstrear the c-mos ATG, also inhibited expression from a heterologous
\*\*\*promoter\*\*\* . In addition to NIH 3T3 cells, the c-mos negative regulatory sequence was active in BALB/3T3 cells, PC12 rat pheochromocytoma cells, and A549 human \*\*\*lung\*\*\* \*\*\*carcinoma\*\* cells. Site- \*\*\*specific\*\*\* mutagenesis identified three possibly interacting regions that were involved in negative regulatory activity, located around -460, -425, and -405 with respect to the ATG. RNase protection analysis indicated that once the negative regulatory sequences were deleted, transcription in NIH 3T3 cells initiated from the same transcription initiation sites normally utilized in spermatocytes, approximately 280 nucleotides upstream of the ATG. Deletions beyond the spermatocyte \*\*\*\*promoter\*\*\* , however, allowed transcription initiation from progressively downstream c-mos sequences. Deletion or mutation of sequences surrounding the ocotyte \*\*\*\*promoter\*\*\* at -55 also had little effect on expression of c-mos constructs in NIH 373 cells. Therefore, the major determinant of c-mos expression in NIH 3T3 cells was removal of the

negative regulatory sequence rather than the utilization of a unique
\*\*\*promoter\*\*\*. The c- mos negative regulatory sequences thus appear to play a significant role in tissue-specific c-mos expression by inhibiting

transcription in somatic cells.

=> d his

#### (FILE 'HOME' ENTERED AT 16:15:29 ON 19 JAN 2005)

FILE 'BIOSIS, EMBASE, CAPLUS' ENTERED AT 16:15:40 ON 19 JAN 2005 1526 S EGF 2 OR EPITHELIAL GLYCOPROTEIN 2 OR EP CAM OR 17 1A OR GA73

52 S L1 AND (PROMOTER OR REGULA? ELEMENT OR REGULAT? REGION OR 5 U

37 DUP REM L2 (15 DUPLICATES REMOVED)

5817 S CARCINOMA (3A) (SELECT? OR RESTRICT? OR SPECIFIC?) 235 S L4 AND (PROMOTER OR REGULA? ELEMENT OR REGULAT? **REGION OR 5** 

6 S L5 AND LUNG CARCINOMA 16

4 DUP REM L6 (2 DUPLICATES REMOVED)

=> s I5 and lung L8 24 L5 AND LUNG

=> dup rem 18

PROCESSING COMPLETED FOR L8

20 DUP REM L8 (4 DUPLICATES REMOVED)

=> s l9 not l7 L10 16 L9 NOT L7

YOU HAVE REQUESTED DATA FROM 16 ANSWERS - CONTINUE? Y/(N):y

L10 ANSWER 1 OF 16 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation, on

AN 2003:106886 BIOSIS

DN PREV200300106886

- The variant hepatocyte nuclear factor 1 activates the P1 \*\*\*promoter\*\*\* of the human alpha-folate receptor gene in ovarian carcinoma.
- AU Tomassetti, Antonella [Reprint Author]; Mangiarotti, Fabio; Mazzi, Mimma; Sforzini, Sabrina: Miotti, Silvia; Galmozzi, Enrico; Elwood, Patrick C.; Canevari, Silvana
- CS Unit of Molecular Therapies, Department of Experimental Oncology, Istituto Nazionale Tumori, Via Venezian 1, 20133, Milan, Italy antonella.tomassetti@istitutotumori.mi.it
- SO Cancer Research, (February 1 2003) Vol. 63, No. 3, pp. 696-704. print. ISSN: 0008-5472 (ISSN print).

DT Article

General Review; (Literature Review)

A English

ED Entered STN: 26 Feb 2003

Last Updated on STN: 26 Feb 2003

AB The alpha folate receptor (alphaFR) is a membrane glycoprotein that binds folates, and mediates their uptake and that of antifolate drugs. alphaFR is absent on ovarian surface epithelium (OSE) but is detectable during early transforming events in this epithelium, with increasing expression levels in association with tumor progression. Analysis of transcriptional regulation of the alphaFR gene have revealed two \*\*\*promoter\*\*\*
regions, P1 and P4, flanking exons 1 and 4, respectively, and a
requirement for three SP1 sites and an INR element for optimal P4 activity. Here, we focused on the P1 transcription regulation in ovarian carcinoma cells. RNase protection assay indicated that the 5'-untranslated region is heterogeneous because of different start sites and alternative splicing of exon 3. A core region of the P1

\*\*\*promoter\*\*\* was sufficient for maximal \*\*\*promoter\*\*\* activity in

ovarian carcinoma cell lines but not in OSE cells or in alphaFR-nonexpressing cell lines. Deletion and mutation analysis of this core \*\*\*promoter\*\*\* identified a cis- \*\*\*regulatory\*\*\*

\*\*\*element\*\*\* at position +27 to +33 of the untranslated exon 1, which is responsible for maximum P1 activity. This element formed an abundant DNA-protein complex with nuclear proteins from ovarian cancer cells but not from other cell lines or OSE cells. Competition experiments and supershift assays demonstrated binding of the P1 cis- \*\*\*regulatory\*\*\*

\*\*\*element\*\*\* by a transcription factor involved in embryonic development, the variant hepatocyte nuclear factor-1 (vHNF1). Analysis of RNA from various cell lines and surgical specimens confirmed that vHNF1 is expressed in ovarian carcinomas. Thus, vHNF1 regulates tissue
\*\*\*specific\*\*\* transcription in ovarian

\*\*\*carcinoma\*\*\*.

L10 ANSWER 2 OF 16 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation, on STN

AN 1999:484497 BIOSIS DN PREV199900484497

- TI DNA vaccination against the ovarian carcinoma-associated antigen folate receptor alpha (FRalpha) induces cytotoxic T lymphocyte and antibody responses in mice.
- AU Neglia, Francesca; Orengo, Anna Maria; Cilli, Michele; Meazza, Raffaella; Tomassetti, Antonella; Canevari, Silvana; Melani, Cecilia; Colombo, Mario P.; Ferrini, Silvano [Reprint author]
- CS Centro di Biotecnologie Avanzate, Istituto Nazionale per la Ricerca sul Cancro, Largo Rosanna Benzi No. 10, 16132, Genova, Italy
- Cancer Gene Therapy, (July-Aug., 1999) Vol. 6, No. 4, pp. 349-357. print. ISSN: 0929-1903.
- DT Article
- LA English ED Entered STN: 16 Nov 1999
- Last Updated on STN: 16 Nov 1999

Last Updated on STN: 16 Nov 1999

AB Human folate receptor alpha (FRalpha) is a folate-binding protein that is 
\*\*\*selectively\*\*\* overexpressed in ovarian 
\*\*\*Carcinoma\*\*\* and has been regarded as a suitable target antigen for immunotherapy purposes. To study the possible use of this antigen in DNA vaccination, FRalpha cDNA was ligated into the VR1012 (Vical) expression vector under the transcriptional control of the cytomegalovirus 
\*\*\*promoter\*\*\*. A total of 100 mug of purified plasmid DNA was injected intramuscularly in 
BALB/c mice three times at 14-day intervals. At 10 days after the second injection, the sera of the animals (100%) displayed significant antibody titers (by indirect immunofluorescence and fluorescence-activated cell 
sorder analysis) against syngneric C28 cells transcluced with FRalpha but sorter analysis) against syngeneic C26 cells transduced with FRalpha, but not against unmodified C26 cells. Immunoglobulin G2a was the predominant isotype. In addition, specific cytotoxic T lymphocyte activity against FRalpha-transduced C26 cells could be detected in splenocytes from all immunized animals. Coinjection of a plasmid containing interleukin-2 cDNA increased both antibody titers and cytotoxic T lymphocyte activity. increased both antibody titers and cytotoxic T lymphocyte activity.

Challenge by subcutaneous injection with FRalpha-transduced C26 cells (performed 10 days after the third injection) showed a statistically significant delay in tumor growth. Vaccination with the FRalpha and interleukin-2 cDNA mixture, which was performed after an intravenous injection of FRalpha-transduced cells, enhanced the mean survival time and reduced the number of \*\*\*Iung\*\*\* metastases, thus suggesting that such vaccination is effective even against preexisting tumor cells.

L10 ANSWER 3 OF 16 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN
AN 2001365405 EMBASE
TI Molecular detection of p16 \*\*\*promoter\*\*\* methylation in the serum of patients with esophageal squamous cell carcinoma.

AU Hibi K.; Taguchi M.; Nakayama H.; Takase T.; Kasai Y.; Ito K.; Akiyama S.;

CS K. Hibi, Second Department of Surgery, Nagoya University School of Medicine, 65 Tsurumai-cho, Showa-ku, Nagoya 466-8550, Japan. khibi@med.nagoya-u.ac.jp SO Clinical Cancer Research, (2001) 7/10 (3135-3138).

Refs: 19 ISSN: 1078-0432 CODEN: CCREF4

CY United States Journal; Article

FS 005 General Pathology and Pathological Anatomy

Surgery

016

Human Genetics 022

Gastroenterology

LA English

AB Purpose and Experimental Design: Recent evidence shows that the presence of \*\*\*promoter\*\*\* hypermethylation of tumor suppressor genes has been demonstrated in the serum DNA of patients with various cancers such as

demonstrated in the serum DNA of patients with various cancers such as 
""lung"", liver, and head and neck cancer. We have examined
""promoter"" hypermethylation of the p16 gene in esophageal squamous cell ""carcinoma" (SCC) using methylation- ""specific" PCR to detect tumor DNA in the serum. Results: Aberrant ""promoter" methylation of the p16 gene was detected in 31 of 38 (82%) esophageal SCCs. Subsequently, we tested for ""promoter" methylation in the paired serum DNA of 31 patients with a p16 alteration in the primary threat. Wile found that 7 of these 31 (23%) price to be the same are thylation. tumor. We found that 7 of these 31 (23%) patients had the same methylation changes in the serum DNA. Conclusions: This result indicates that

\*\*\*promoter\*\*\* methylation present in the tumors of esophageal SCC

patients can be detected in the serum of the same patient and that this approach can potentially be used for the screening and monitoring of the

L10 ANSWER 4 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2004:930659 CAPLUS

TI Cancer-specific activation of the survivin \*\*\*promoter\*\*\* and its

potential use in gene therapy J. Chen, Jin-Shing, Liu, Jaw-Ching, Shen, Lei, Rau, Kung-Ming, Kuo, Hsu-Ping,

Li, Yan M.; Shi, Daren; Lee, Yung-Chie; Chang, King-Jen; Hung, Mien-Chie. 5 Department of Molecular and Cellular Oncology, The University of Texas MD

Anderson Cancer Center, Houston, TX, 77030, USA SO Cancer Gene Therapy (2004), 11(11), 740-747 CODEN: CGTHEG; ISSN: 0929-1903

PB Nature Publishing Group

Journal

English AB Survivin is expressed in many cancers but not in normal adult tissues and is transcriptionally regulated. To test the feasibility of using the is transcriptionally regulated. To test the feasibility of using the survivin \*\*\*promoter\*\*\* to induce cancer-specific transgene expression in \*\*\*lung\*\*\* cancer gene therapy, a vector expressing a luciferase gene driven by the survivin \*\*\*promoter\*\*\* was constructed and evaluated in vitro and in vivo. It was found that the survivin \*\*\*promoter\*\*\* was generally more highly activated in cancer cell lines than in normal and immortalized normal cell lines. When delivered i.v. by DNA:liposome complexes, the survivin \*\*\*promoter\*\*\* was more than 200 times more cancer specific than the cytomegalovirus \*\*\*promoter\*\*\* in vivo. To identify \*\*\*lung\*\*\* cancer patients who may benefit from gene therapy with the survivin \*\*\*promoter\*\*\*, survivin protein expression was measured in surgical specimens of 75 non-small-cell \*\*\*lung\*\*\* cancers and 10 normal \*\*\*lung\*\*\* tissues by immunohistochem, staining and found that survivin is expressed in most of the non-small-cell \*\*\*lung\*\*\* cancers tested (81%, 61 of 75) but none of the normal \*\*\*lung\*\*\* tissues. The survivin \*\*\*promoter\*\*\*, survivin is expressed the growth of cancer cells in vitro and in vivo. These results suppressed the growth of cancer cells in vitro and in vivo. These results indicate that the survivin \*\*\*promoter\*\*\* is a cancer-specific \*\*\*promoter\*\*\* for various cancers and that it may be useful in cancer

gene therapy.

RE.CNT 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L10 ANSWER 5 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN

AN 2003:413860 CAPLUS

DN 139:917

Dual specificity tumor killing viral vectors driven by the telomerase

\*\*\*promoter\*\*\* and uses for cancer gene therapy
Irving, John M.; Karpf, David B.; Schiff, J. Michael

USA

SO U.S. Pat. Appl. Publ., 25 pp.

CODEN: USXXCO

DT Patent LA English

FAN.CNT 1 PATENT NO.

APPLICATION NO.

DATE 20020725

PI US 2003099616 A1 20030529 US 2002-206447 PRAI US 2001-308029P P 20010725

KIND DATE

AB The present invention discloses the specificity of multiple reascriptional regulatory elements can be combined to make adenoviral vector systems that selectively target cancer cells and its uses in gene therapy for cancers. The \*\*\*promoter\*\*\* for telomerase reverse transcriptase (TERT) can be combined in a remarkably synergistic fashion with another \*\*\*promoter\*\*\* that has expression restricted to cancer cells or a particular tissue type. The two promoters work synergistically for exquisite targeting of the malignant cells-where it causes cell lysis while leaving neighboring healthy cells intact. This invention also includes methods for constructing and selecting the viral vectors, host cells transduced with the vector construct, and the host cells monitored for any effect of the vector.

L10 ANSWER 6 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2003:178335 CAPLUS DN 138:231393

Tumor-specific gene therapy for undifferentiated thyroid carcinoma using the human telomerase reverse transcriptase AU Takeda, Teiji; Hashizume, Kiyoshi \*\*promoter\*

CS Department of Aging Medicine and Geriatrics, Shinshu University School of Medicine, Japan

SO Horumon to Rinsho (2003), 51(2), 149-154 CODEN: HORIAE; ISSN: 0045-7167

PB Igaku no Sekaisha DT Journal

Japanese

Japanese

3 The authors previously developed recombinant adenoviruses carrying herpes simplex virus thymidine kinase (HSVtk) genes to evaluate the possibility of tissue-specific gene therapy for thyroid carcinoma. The HSVtk gene was driven by a minimal thyroglobulin (TG) \*\*\*promoter\*\*\* (AdTGtk) and a tandemly repeated minimal TG \*\*\*promoter\*\*\* (Ad2.times.TGtk) to obtain thyroid-specific cell killing ability. Ad2.times.TGtk showed a beneficial effect for tissue-specific gene therapy for TG-producing thyroid carcinoma, but not for undifferentiated thyroid carcinoma. The authors

placed HSVtk gene under the control of human telomerase reverse transcriptase (hTERT) gene ""promoter" (AdhTERTtk). Tumor-specific transcriptional activity by hTERT """promoter" was confirmed. The transduction of HSVtk genes by infection with AdhTERTtk followed by ganciclovir (GCV) treatment showed powerful cytotoxicity for TG-producing and non-TG-producing thyroid carcinoma cell lines but no or little cytotoxicity for normal cell lines. After adenovirus/GCV treatment for ARO tumor-bearing nude mice, AdhTERTtk inhibited the tumor growth. Ad2.times.TGtk/GCV and AdhTERTtk/GCV treatment showed no or very little cytotoxicity in liver, kidney, spleen, thyroid, \*\*\*lung\*\*\*, and testis. These data suggest a beneficial effect of AdhTERTtk for gene therapy of undifferentiated thyroid carcinoma without toxicity for normal

L10 ANSWER 7 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2002:897718 CAPLUS

DN 138:120721

- TI Characterization of a tissue-specific CDP/Cux isoform, p75, activated in breast tumor cells
- AU Goulet, Brigitte; Watson, Peter; Poirier, Madeleine; Leduy, Lam; Berube, Ginette; Meterissian, Sarkis; Jolicoeur, Paul; Nepveu, Alain CS Molecular Oncology Group, McGill University Health Center, Montreal, QC,
- H3A 1A1, Can.
- SO Cancer Research (2002), 62(22), 6625-6633 CODEN: CNREA8; ISSN: 0008-5472
- PB American Association for Cancer Research

DT Journal

LA English

AB Two isoforms of the CCAAT-displacement protein/cut homeobox (CDP/Cux) transcription factor have been characterized thus far. The full length protein, p200, which contains four DNA binding domains, transiently binds to DNA and carries the CCAAT-displacement activity. The p110 isoform is generated by proteolytic processing at the G1-S transition and is capable of stable interaction with DNA. Here the authors demonstrate the existence of a shorter CDP/Cux isoform, p75, which contains only two DNA binding domains, Cut repeat 3 and the Cut homeodomain, and binds more stably to DNA. CDP/Cux p75 was able to repress a reporter carrying the ""promoter" for the cyclin-dependent kinase inhibitor p21 gene and to activate a DNA polymerase alpha, gene reporter. Expression of CDP/Cux

p75 involved a novel mechanism: transcription initiation within intron 20. The intron 20-initiated mRNA (I20-mRNA) was expressed at higher level in the thymus and in CD4+/CD8+ and CD4+ T cells. I20-mRNA was expressed only

weakly or not at all in normal human mammary epithelial cells and normal breast tissues but was detected in many breast tumor cells lines and breast tumors. In invasive tumors a significant assocn, was established between higher I20-mRNA expression and a diffuse infiltrative growth pattern. In agreement with these findings, T47D breast cancer cells stably expressing p75 could not form tubule structures in collagen but rather developed as solid undifferentiated aggregates of cells. Taken together, these results suggest that aberrant expression of the CDP/Cux p75 isoform in mammary epithelial cells may be assocd. with the process of

tumorigenesis in breast cancer.
RE.CNT 72 THERE ARE 72 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L10 ANSWER 8 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2002:393088 CAPLUS

- Tumour specific \*\*\*promoter\*\*\* region methylation of the human homologue of the Drosophila Roundabout gene DUTT1 (ROBO1) in human
- AU Dallol, Ashraf, Forgacs, Eva; Martinez, Alonso; Sekido, Yoshitaka; Walker, Rosemary, Kishida, Takeshi, Rabbitts, Pamela; Maher, Eamonn R.; Minna, John D.; Latif, Farida
- CS Section of Medical and Molecular Genetics, Department of Paediatrics and Child Health, The Medical School, University of Birmingham, Birmingham, B15 2TT, UK
- SO Oncogene (2002), 21(19), 3020-3028 CODEN: ONCNES; ISSN: 0950-9232 PB Nature Publishing Group

DT Journal

LA English
AB The human homolog of the Drosophila Roundabout gene DUTT1 (Deleted in

Twenty Twenty) or ROBO1 (Locus Link ID 6091), a member of the NCAM family Iwenty Iwenty of ROBO1 (Locus Link to 0031), a member of the NOAM family of receptors, was recently cloned from the \*\*\*\*lung\*\*\* cancer tumor suppressor gene region 2 (LCTSGR2 or U2020 region) at 3p12. DUTT1 maps within a region of overlapping homozygous deletions characterized in both small cell \*\*\*\*lung\*\*\* cancer lines (SCLC) and in a breast cancer line. In this report the authors (a) defined the genomic organization of the DUTT1 gene, (b) performed mutation and expression anal. of DUTT1 in \*\*\*lung\*\*\* , breast and kidney cancers, (c) identified tumor specific
\*\*\*promoter\*\*\* region methylation of DUTT1 in human cancers. The gene was found to contain 29 exons and spans at least 240 kb of genomic sequence. The 5' region contains a CpG island, and the poly(A)+ tail has an atypical 5'-GATAAA-3' signal. The authors analyzed DUTT1 for mutations in \*\*\*lung\*\*\*\*, breast and kidney cancers; no inactivating mutations were detected by PCR-SSCP. However, seven germline missense changes

found and characterized. DUTT1 expression was not detectable in one out of 18 breast tumor lines analyzed by RT-PCR. Bisulfite sequencing of the

```
***promoter*** region of DUTT1 gene in the HTB-19 breast tumor cell line
   (not expressing DUT11) showed complete hypermethylation of CpG sites within the ""promoter"" region of the DUTT1 gene (-244 to +27 relative to the translation start site). The expression of DUTT1 gene was reactivated in HTB-19 after treatment with the demethylating agent
    5-aza-2'-deoxycytidine. The same region was also hypermethylated in six
   out of 32 (19%) primary invasive breast carcinomas and eight out of 44 (18%) primary clear cell renal cell carcinomas (CC-RCC) and in one out of
     26 (4%) primary NSCLC tumors. Furthermore 80% of breast and 75% of CC-
    tumors showing DUTT1 methylation had allelic losses for 3p12 markers hence
   obeying Knudson's two hit hypothesis. The authors' findings suggest that DUTT1 warrants further anal. as a candidate for the tumor suppressor gene
    (TSG) at 3p12, a region defined by hemi and homozygous deletions and
    functional anal.
RE.CNT 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS
RECORD
           ALL CITATIONS AVAILABLE IN THE RE FORMAT
L10 ANSWER 9 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2001:654737 CAPLUS
      135:206499
```

TI Non-squamous epithelium-specific EGP-2 \*\*\*promoter\*\*\* driven transcription for cancer therapy De Leij, Lou Franciscus Maria Hubertus; McLaughlin, Pamela Marijke Jane; Ruiters, Marcel Herman Josef; Harmsen, Martin Conrad; Van der Molen, Henk; Terpstra, Peter; Dokter, Willem Hendrik Abraham Rijksuniversiteit te Groningen, Neth. SO Eur. Pat. Appl., 21 pp. CODEN: EPXXDW DT Patent LA English FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE P 1130106 A1 20010905 EP 2000-200728 20000301 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO PI EP 1130106

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
CA 2364314 AA 20010927 CA 2001-2364314 20010228
WO 2001071015 A3 20020131
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
EP 1190085 A2 20020327 EP 2001-952047 20010228
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
NZ 515206 A 20040326 NZ 2001-515206 20010228
US 2002156041 A1 20021024 US 2002-9579 20020326
PRAI EP 2000-200728 A 20000301
WO 2001-NL166 W 20010228
AB The invention relates to the field of cancer therapy and diagnosis, in

AB The invention relates to the field of cancer therapy and diagnosis, in particular of carcinomas. The invention provides an isolated and/or recombinant nucleic acid comprising a tissue specific \*\*\*promoter or functional fragment thereof allowing for expression of a nucleic acid of interest operably linked to said \*\*\*promoter\*\*\* or functional fragment thereof in a cancer cell wherein said expression in said cancer cell is essentially \*\*\*carcinoma\*\*\* \*\*\*selective\*\*\* . In a preferred embodiment, the invention provides the isolation and use of EGP-2 transcriptional regulatory sequences to regulate transient expression of the cytosine dearninase gene in EGP-2 expressing carcinoma cells. The invention further provides a vector or gene delivery vehicle comprising a nucleic acid according to the invention. Such gene delivery vehicles as provided by the invention are very useful in carcinoma

therapy, or in therapy directed at non-squamous epithelium.
RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L10 ANSWER 10 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2001:380649 CAPLUS

TI Antigen-binding fragments specific for dendritic cells, compositions and methods of use thereof antigens recognized thereby and cells obtained

Schmitz, Juergen; Dzionek, Andrzej; Buck, David William Miltenyi Biotech G.m.b.H., Germany

SO PCT Int. Appl., 114 pp. CODEN: PIXXD2

DT Patent

LA English FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2001036487 20010525 WO 2000-IB1832 20001115 WO 2001036487 АЗ 20020510 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,

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CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
CA 2396428 AA 20010525 CA 2000-2396428 20001115
EP 1301539 A2 20030416 EP 2000-979855 20001115
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR
                        IE, FI, CY, TR
 JP 2004512006 T2
PRAI US 1999-165555P
                                                                         20040422 JP 2001-538976
P 19991115
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         US 1999-167076P
US 2000-179003P
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                                                                               19991123
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          US 2000-180775P
                                                                                20000207
         US 2000-196824P
US 2000-197205P
                                                                               20000411 20000413
         WO 2000-IB1832
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                                                                               20001115
 AB The invention provides antigen-binding fragments specific for dendritic cells and effective in treatment and/or diagnosing a variety of disorders. Methods of use are also provided as are methods for screening for addnl.
          such antigen-binding fragments and the products obtained thereby
 L10 ANSWER 11 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2001;310088 CAPLUS
 TI p16liNK4a and histology-specific methylation of CpG islands by exposure to tobacco smoke in non-small cell ***lung*** cancer
AU Kim, Duk-Hwan; Nelson, Heather H.; Wiencke, John K.; Zheng, Shichun;
 Christiani, David C.; Wain, John C.; Mark, Eugene J.; Kelsey, Karl T.
CS Department of Environmental Health, Harvard School of Public Health,
 Boston, MA, 02115, USA
SO Cancer Research (2001), 61(8), 3419-3424
CODEN: CNREA8; ISSN: 0008-5472
  PB American Association for Cancer Research
           Journal
LA English

AB The p16INK4a protein inhibits cyclin-dependent kinase 4, a key regulator of progression through the G1 phase of the cell cycle. Methylation of CpG islands in the ""promoter" region is an important avenue for inactivation of p16. The mechanism of methylation of the p16 ""promoter" region, however, has not been elucidated. Recent reports investigating p16 methylation in non-small cell ""lung" cancer (NSCLC) suggest that carcinogens in tobacco smoke induce the DNA methylation process. We investigated the assocn, between methylation of the p16 ""promoter" region and exposure to tobacco smoke in 185 primary NSCLCs. We also studied the relationship of p16 methylation with multiling of the K-rss and p53 genes as well as with methylation at the
         mutation of the K-ras and p53 genes, as well as with methylation at the 
DAP-kinase and p14ARF loci. Finally, we evaluated the prognostic 
significance of p16 methylation in NSCLC. The prevalence of p16
       significance of p16 methylation in NSCLC. The prevalence of p16 methylation was greater in squamous cell carcinoma (41%) compared with adenocarcinoma (22%; P = 0.03; Fisher's exact test). Methylation of p16 was significantly assocd, with pack-years smoked (P = 0.007; Wilcoxon rank sum test), duration of smoking (P = 0.0009; Wilcoxon rank sum test), and neg, with the time since quitting smoking (P = 0.03; Wilcoxon rank sum test). No methylation of the nearby p14ARF locus was detected, and methylation of the DAP-kinase locus was not assocd, with either p16
        methylation or with exposure to tobacco smoke. In patients with stage 1
        adenocarcinoma, p16 methylation was an independent risk factor predicting significantly shorter postsurgery survival (P = 0.03), controlling for the
        significant effects of other factors, including K-ras mutation. These findings suggest that methylation of CpG islands in tobacco-assocd. cancers occurs in a gene- and tissue-specific manner and is induced
 directly or indirectly by exposure to tobacco smoke in NSCLC.
RE.CNT 53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS
RECORD
                        ALL CITATIONS AVAILABLE IN THE RE FORMAT
L10 ANSWER 12 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 2001:247215 CAPLUS
 DN 134:276498
TI Engineering of replication selective adenoviruses with tumor-associated antigen ***promoter*** for use in cancer therapy
         Molnar-kimber, Katherine; Toyoizumi, Takane
           The Trustees of the University of Pennsylvania, USA
        PCT Int. Appl., 56 pp.
CODEN: PIXXD2
DT Patent
 LA English
FAN CNT 1
        PATENT NO.
                                                            KIND DATE
                                                                                                           APPLICATION NO.
         WO 2001023004
                                                                    A1 20010405 WO 2000-US27212
                                                                                                                                                                                  20001002
             WO 2001023004 A1 20010405 WO 2000-US27212 20001002
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
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in cancers and malignancies, as well as in proliferative cells, characterizing diseases, such as restenosis, intimal proliferative disease
       and pulmonary hypertension. The selected Ad vectors are driven by promoters of the tumor assocd, antigens, or RNA transcripts or genes therefor, substituting for the activity of at least adenovirus E1A
       ***promoter***, which has been deactivated or diminished. Also provided is the use of the Ad vector to deliver therapeutic compns. to patients, as
       well as a method for treating cancers, such as CEA pos. cancers, or
       proliferative cell diseases in a patient by administering to the patient an effective amt. of the Ad vector, which may also express a therapeutic
       gene or peptide, and treatment may also be combined with radiation, chemotherapy or immunomodulatory agents. The Ad is designed to replicate within the tumor cell, thereby spreading throughout the tumor nodule.
       This permits the delivery of a much higher dose of the heterologous therapeutic protein than previously possible, and the virus achieves a
       direct, oncolytic effect on the tumor.

CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
 RECORD
                    ALL CITATIONS AVAILABLE IN THE RE FORMAT
 L10 ANSWER 13 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN
          1999:582108 CAPLUS
132:120711
DN 132:120/11
TI Tumour-specific arginine vasopressin ***promoter*** activation in small-cell ***lung*** cancer
AU Coulson, J. M.; Stanley, J.; Woll, P. J.
CS CRC Department of Clinical Oncology, University of Nottingham, NG5 1PB, UK
SO British Journal of Cancer (1999), 80(12), 1935-1944
CODEN: BJCAAI; ISSN: 0007-0920
PB Churchill Livingstone
         Journal
          English
 AB Small-cell ***lung*** cancer (SCLC) can produce numerous mitogenic
       neuropeptides, which are not found in normal respiratory epithelium.
Arginine vasopressin is detected in up to two-thirds of SCLC tumors whereas normal physiol. expression is essentially restricted to the
      whereas norms. This presents the opportunity to identify elements of the gene ***promoter*** which could be exploited for SCLC-specific targeting. A series of human vasopressin 5' ***promoter*** fragments (1048 bp, 468 bp and 199 bp) were isolated and cloned upstream of a reporter gene. These were transfected into a panel of ten cell lines,
       including SCLC with high or low endogenous vasopressin transcription, non-SCLC and bronchial epithelium. All these fragments directed reporter gene expression in the five SCLC cell lines, but had negligible activity
      in the control lines. The level of reporter gene expression reflected the level of endogenous vasopressin prodn., with up to 4.9-fold (s.d. 0.34) higher activity than an SV40 ***promoter***. The elements required for this strong, restricted, SCLC-specific **promoter*** activity are contained within the 199-bp fragment. Further anal. of this region indicated involvement of E-box transcription factor binding sites,
although tumor-specificity was retained by a 65-bp minimal
""promoter"" fragment. These data show that a short region of the
vasopressin ""promoter"" will drive strong expression in SCLC in
vitro and raise the possibility of targeting gene therapy to these tumors.

RECNT 37 THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS
 RECORD
                  ALL CITATIONS AVAILABLE IN THE RE FORMAT
L10 ANSWER 14 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN
          1999:244743 CAPLUS
 DN 130:276738
 TI Inducing tumor-specific cytotoxicity using vectors containing H19 or
insulin-like growth factor gene regulatory elements

IN Hochberg, Abraham; Ayesh, Suhail

PA Yissum Research Development Company of the Hebrew University of
Jerusalem,
      Israel
SO PCT Int. Appl., 67 pp.
CODEN: PIXXD2
DT Patent
 LA English
FAN CNT 4
       PATENT NO.
                                                  KIND DATE
                                                                                         APPLICATION NO.
                                                                                                                                              DATE
PI WO 9918195
                                                    A2 19990415 WO 1998-IL486
                                                                                                                                           19981004
          VO 9918195 A3 19990812

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
A 2308124 AA 19990415 CA 1998-2308124 19981004
U 9894571 A1 19990427 AU 1998-94571 19981004
                                                           19990812
       CA 2308124
       AU 9894571
                                               B2 20021219
A2 20000719
       AU 755774
       EP 1019499
                                                          20000719 EP 1998-947759
                                                                                                                                        19981004
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CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-157224P P 19990930
AB The invention provides a replication selective adenovirus (Ad) mutant with improved selectively for tumor cells expressing the tumor assocd. antigen

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IE, FI
BR 9812717
                                                                                                                                                   US 2001011128 A1 20010802 US 2000-739223
PRAI US 1996-26678P P 19960925
WO 1997-CA691 W 19970922
                                         20000822 BR 1998-12717
                                                                                              19981004
                                  T2 20011023 JP 2000-514993
C2 20031020 RU 2000-111553
A 20000602 NO 2000-1684
      JP 2001519148
                                                                                                 19981004
     RU 2214280
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     NO 2000001684
                                                                                                 20000331
                                                                                                                                                        US 1999-276005
                                        A 1997 .
7 19981004
Theolfic
                                                                                                                                                   AB The present invention relates to a tumor-specific ***promoter***, the Hex II ***promoter***, for use in gene targeted therapy that is
PRAI US 1997-943608
WO 1998-IL486
                                                19971003
                                     w
                                                                                                                                                        differentially regulated in cancer cells. The present invention also relates to a gene construct, which includes the Hex II ***promoter*** in a vector selected from pCAT basic expression vector p.DELTA.ElsplB,
 AB The invention relates to the specific expression of heterologous
     sequences, particularly genes encoding cytotoxic products, in tumor cells under the control of regulatory transcriptional sequences. Particularly
     preferred promoters include H19 regulatory sequences, the IGF-1

***promoter***, and the IGF-2 P3 and P4 promoters from genomically imprinted genes. The invention provides expression constructs and methods of administering such expression constructs. The H19 regulatory sequences facilitate expression of a heterologous gene in five different bladder cancer cell lines (HT-1376, EJ28, T24P, 1197, and UM-UC-3). When
                                                                                                                                                       called pHexII4557-CAT, and a shuttle plasmid which includes either .beta-gal or HSV Tk, called p.DELTA.ElspIBHex-LacZ and p.DELTA.ElspIBHex-TK.
                                                                                                                                                   REICHT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
                                                                                                                                                   RECORD
                                                                                                                                                                 ALL CITATIONS AVAILABLE IN THE RE FORMAT
     transfected into bladder cancer cell, an H19/HSV-TK expression plasmid induces bladder cancer cell-specific cytotoxicity in the presence of
     ganciclovir. The compns. and methods of the invention are useful in the
     treatment of cancer.
                                                                                                                                                   ---Logging off of STN---
L10 ANSWER 15 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 1998;793064 CAPLUS
 DN 130:35133

    Pselectin translocation to vascular epithelial lumen by ionizing radiation, and therapeutic use
    Hallahan, Dennis E.; Virudachalam, Subbulakshmi
    Arch Development Corporation, USA
    O PCT Int. Appl., 178 pp.
    CODEN: PIXXD2

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    WO 9853852
A1 19981203 WO 1998-US10913 19980529
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CA 2290563
A4 19981203 CA 1998-2290563 19980529
AU 9886570
A1 19981230 AU 1998-86570 19980529
EP 986401
B1 20040225
                                     A1 19981203 WO 1998-US10913
PI WO 9853852
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PRAI US 1997-48141P
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     WO 1998-US10913
 AB The present invention relates to the use of P-selectin as a targeting
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     agent in radiotherapies for vascular related disease. P-selectin is
                                                                                                                                                   NEWS 1 Web Page URLs for STN Seminar Schedule - N. America
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STN Express with Discover!
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NEWS 5 NOV 30 PHAR reloaded with additional data
     translocated to the lumen of vascular endothelia as a result of radiation
     Thus, P-selecting provides a target for receptor-mediated delivery of drugs, including anticancer drugs and drugs for the treatment of vascular
     disease. However, P-selectin also plays a role in the activation of
     certain inflammatory cells and, as such, plays a role in radiation-induced inflammation. By interfering with P-selectin induction of inflammation, it is possible to modulate inflammatory responses to radiation therapy.
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NEWS 7 DEC 09 12 databases to be removed from STN on December 31, 2004
RE.CNT 6
                                                                                                                                                    NEWS 8 DEC 15 MEDLINE update schedule for December 2004
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                                                                                                                                   NEWS 9 DEC 17 ELCOM reloaded; updating to resume; current-awareness
                                                                                                                                                                    alerts (SDIs) affected
L10 ANSWER 16 OF 16 CAPLUS COPYRIGHT 2005 ACS on STN AN 1998:210872 CAPLUS
                                                                                                                                                   NEWS 10 DEC 17 COMPUAB reloaded; updating to resume; current-awareness
                                                                                                                                                   alerts (SDIs) affected
NEWS 11 DEC 17 SOLIDSTATE reloaded, updating to resume; current-
DN 128:266956
TI Hex II tumor-specific ***promoter*** and its use in gene-targeted
                                                                                                                                                                   alerts (SDIs) affected
     cancer therapy
     Batist, Gerald; Katabi, Maha
                                                                                                                                                   NEWS 12 DEC 17 CERAB reloaded; updating to resume; current-awareness
PA McGill University, Can.; Batist, Gerald; Katabi, Maha
SO PCT Int. Appl., 20 pp.
CODEN: PIXXD2
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NEWS 13 DEC 17 THREE NEW FIELDS ADDED TO IFIPAT/IFIUDB/IFICDB
                                                                                                                                                   NEWS 14 DEC 30 EPFULL: New patent full text database to be available on STN NEWS 15 DEC 30 CAPLUS - PATENT COVERAGE EXPANDED
DT Patent
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FAN CNT 2
                                                                                                                                                   February 2005
NEWS 17 JAN 11 CA/CAPLUS - Expanded patent coverage to include Russia
     PATENT NO.
                                  KIND DATE
                                                              APPLICATION NO.
                                                                                                   DATE
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                                     A1 19980402 WO 1997-CA691
                                                                                                   19970922
        VO 9813507 A1 19980402 WO 1997-CA691 19970922
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RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GP, ET, TLI MR, NI, PT, SE, BE, EL, CE, CG, CL, CM, GA
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           W. B., R., E., IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
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9742927 A1 19980417 AU 1997-42927 19970922
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EP 954590

19991110 EP 1997-918865

19970922

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L1 248 LUNG CARCINOMA (3A)(SELECT? OR SPECIFIC? OR RESTRIC?)

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ELEMENT OR 5 UTR)

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=> dup rem 12 PROCESSING COMPLETED FOR L2 L3 2 DUP REM L2 (2 DUPLICATES REMOVED)

L3 ANSWER 1 OF 2 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation.

**DUPLICATE 1** 

AN 2001:225975 BIOSIS DN PREV200100225975

TI Adenovirus-mediated suicide gene transfer to small cell \*\*\*lung\*\*\*
\*\*\*carcinoma\*\*\* using a tumor- \*\*\*specific\*\*\* \*\*\*promoter\*\*\*.

AU Morimoto, Emiko; Inase, Naohiko [Reprint author]; Miyake, Shuji;

Yoshizawa, Yasuyuki

CS Pulmonary Medicine, Tokyo Medical and Dental University, 1-5-45 Yushima, Bunkyo-ku, Tokyo, 113-8519, Japan ninase.pulm@md.ac.jp SO Anticancer Research, (January-February, 2001) Vol. 21, No. 1A, pp.

CODEN: ANTRD4. ISSN: 0250-7005. DT Article

LA English

ED Entered STN: 9 May 2001 Last Updated on STN: 18 Feb 2002

AB The gastrin-releasing peptide (GRP) is expressed in most types of small cell lung carcinoma (SCLC) and the GRP \*\*\*promoter\*\*\* is thought to be potentially useful for tumor-specific expression of the suicide gene in potentially useful for tumor-specific expression of the suicide gene in SCLC. We constructed an adenovirus containing the herpes simplex thymidine kinase suicide gene driven by the GRP \*\*\*promoter\*\*\*

(AdGRP-TK) and transfected it into GRP-expressing SCLC cells (SBC5) to confer sensitivity to ganciclovir (GCV). After infection with AdGRP-TK, SBC5 cells became more sensitive to GCV in vitro. In nude mice, a subcutaneously-inoculated tumor of SBC5 cells infected with AdGRP-TK in advance regressed completely after intraperitoneal administration of GCV. These results suggest that adenovirus-mediated gene transfer of the suicide gene followed by pro-drug treatment may be applicable to SCLC.

=> d bib abs 2

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on STN

AN 92191536 EMBASE

DN 1992191536

TI Identification of a negative \*\*\*regulatory\*\* \*\*\*element\*\*\* that inhibits c-mos transcription in somatic cells.

AU Zinkel S.S.; Pal S.K.; Szeberenyi J.; Cooper G.M.
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SO Molecular and Cellular Biology, (1992) 12/5 (2029-2036). ISSN: 0270-7306 CODEN: MCEBD4

United States

DT Journal; Article FS 004 Microbiology

English

SL

English
We have used transient expression assays to identify a cis-acting region in the 5 flanking sequence of murine c-mos which, when deleted, allows expression from the c-mos \*\*\*promoter\*\*\* in NIH 3T3 cells. This negative regulatory sequence, located 400 to 500 nucleotides upstream of the c-mos ATG, also inhibited expression from a heterologous
\*\*\*promoter\*\*\* . In addition to NIH 3T3 cells, the c-mos negative

regulatory sequence was active in BALB/3T3 cells, PC12 rat pheochromocytoma cells, and A549 human \*\*\*lung\*\*\* \*\*\*carcinoma\*\*\*cells. Site- \*\*\*respecific\*\*\* mutagenesis identified three possibly interacting regions that were involved in negative regulatory activity, located around -460, -425, and -405 with respect to the ATG. RNase protection analysis indicated that once the negative regulatory sequences were deleted, transcription in NIH 3T3 cells initiated from the same transcription initiation sites normally utilized in spermatocytes, transcription intribution sites normally utilized in spermatocytes, approximately 280 nucleotides upstream of the ATG. Deletions beyond the spermatocyte \*\*\*promoter\*\*\*, however, allowed transcription initiation from progressively downstream c-mos sequences. Deletion or mutation of sequences surrounding the occyte \*\*\*promoter\*\*\* at -53 also had little effect on expression of c-mos constructs in NIH 373 cells. Therefore, the major determinant of c-mos expression in NIH 3T3 cells was removal of the negative regulatory sequence rather than the utilization of a unique
\*\*\*promoter\*\*\* . The c- mos negative regulatory sequences thus appear to

\*promoter\*\*\* play a significant role in tissue-specific c-mos expression by inhibiting

transcription in somatic cells.

---Logging off of STN---

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS

SINCE FILE TOTAL

**ENTRY** SESSION

**FULL ESTIMATED COST** 

STN INTERNATIONAL LOGOFF AT 17:12:01 ON 19 JAN 2005

---Logging off of STN---

END

Unable to generate the STN prompt. Exiting the script...